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FIME COVERS 1961 - 8 Jul 1891 | VOL 1:7 ISS 1 FIME LAST UPPATED: 8 Jul 1891 | (2002)735/ED

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified offective December 16, 2001. Please theck your CDI profiles to see if they need to be revised. For information or CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

⇒ d all tot 179.

139 ANSWER 1 OF 34 HOAFLUS COPYRIGHT 200. ACS

AN 1000:40:8:8 HCAPLUD

DN: 136:382505

T1 Device for monitoring cells

III Pither, C. Bruce; Hemperly, John Jacob; Guarino, Richard D.; Wodnicka, Magdalena; Stitt, David T.; Burrell, Gregory J.; Foley, Tinothy G., Jr.; Beaty, Patrick Shawn

FA Becton, Dickinson and Company, USA

30 U.S., 41 pp., Cont.-in-part of U.S. Ser. No. 715,120. CODEN: USEMAM

DG Patent

LA English

IC ICM C13Q001-18

MCI 435032000

OC 9-1 (Baschemical Methods) Section pross-reterence(a : 1, 4

FAN. CUT 3

rm.	PATEET NO.	KIND)	LATE	AFFLICATION NO.	DATE
ΡI	US 6395506	Bl	200.08.19	78 1999-342720	199:0629
	EP 509791 EP 509791	Al El	199.10.1 19960703	ME 1392-305/91	199.0415
	E: DE, FE, CA 2060329	AA	199./1019	CA 1992-2060829	199.:0416
	JP 35137596 JP 37673510	A.1 B4	19970601 19950839	JB 1992-99308	19910413
PKAI	US 1991-087359 US 1993-25899	Вì А.:	19910418 19930808		
	US 1096-115537	A	19900918		

AB The present invention relates to methods for detection and evaluation of metabolic activity of eukaryotic and/or prodaryotic cells based upon their ability to consume dissolved exygen. The methods utilize a luminescence detection system which makes use of the sensitivity

ST

ΙΤ

ΙT

ΙT

(device for monitoring cells)

```
of the luminescent emission of certain compds. to the presence
of oxygen, which quenches (diminishes) the compd.'s luminescent
emission in a concn. dependent manner. Respiring eukaryotic
and/or prokaryotic cells will affect the oxygen concn. of a liq.
medium in which they are immersed. Thus, this invention provides a
convenient system to gather information on the presence, identification,
quaritification and cytotexic activity of eukaryotic and/or prokaryotic
wells by deta, their effect on the oxygen concn. of the media in
which they are present.
device monitoring cell,
Plater
   (Microtitration; device for monitoring cells)
Analytical apparatus
Antibiotics
Bacteria (Eubacteria)
Biological materials
B1000
Blood serum
Tell
Fell proliferation
Chemicals
Westing materials
Composition
  Concentration (condition)
Julture media
Cytotoxicity
Drugs
Escherichia coli
Eukaryota
Extracellular matrix
  Fluorescence quenching
Impermeability
Insecta
  Light
Liquids
  Luminescence
  Luminescence quenching
  Luminescence spectroscopy
  Luminescent substances
 Mathematical methods
Metabiliam.
Microsrganism
Molecules
Farticles
Permeability
Proharyote
Iseudimonas aeruginosa
Fadiation
Feducing agents
Festiration, animal
Fespiration, microbial
  Sensors
Colutes
 Wavelength
Wetting
Yeast
   (device for monitoring cells)
FL: ADV (Adverse effect, including toxicity); BIOL (Biological study)
   -device for monitoring cells)
FL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
```

```
Flastics, analysis
ΙΤ
    FL: AFU (Analytical role, unclassified); ANST (Analytical study)
        device for monitoring cells.
    Fubber, analysis
TΤ
    FL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (device for monitoring cells)
    Silicone rubber, analysis
TΤ
    FL: ARU (Analymical role, unclassified); ANST (Analytical study)
        (device for monitoring cells:
ΙΤ
    Growth factors, animal
    FL: BSW (Biological study, unclassified); BIOL Biological study)
        -device for monitoring cells;
    Collagens, billogical studies
ΤT
    FL: BUW (Biological use, unclassified); BI/L (Biological study; USES
     (Us.s)
        (device for monitoring cells)
ΙT
    Entactin
    FL: BUU (Biological use, unclassified); BILL (Biological Study); USES
     (Uses)
        (device for monitoring cells)
    Laminins
TT
    FL: BUU (Biological use, unclassified); BIUL (Biological study); USES
        (device for monitoring cells)
    Proteoglycans, biological studies
TΤ
    FL: BUU (Biological use, unplassified); BELL (Biological study.; USES
        (netaritin sulfate-conty.; device for minitoring cells)
IΤ
    Optical detectors
        (luminescence; device for monitoring cells)
ΙT
    Amimal dell
        (mammal; device for monitoring cells)
    Amino acids, brological studies
ΙT
    FL: BUU (Biological use, unclassified); BIOL (Biological study.; USES
     (Usus)
        inchessential; device for monitoring cells)
ΙT
    Colladens, biological studies
    FL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Usws)
        (type IV; newice for monitoring cells)
    1499-16-1, 3,19-Diphenylanthracene 18188-62-CD, Tris-2,2'-
ΙT
    bapyridylruthenium (II), salts 36309-86-8, Tris-4,7-diphenyl-1,10-
    phenanthroline ruthenium (II) chloride 55525-27-4, Tris-2,2'-
    hapyridylruthenium (II) chloride hexahydrate. 63373-04-6D,
    Tris-4,7-diphenyl-1,11-phenanthroline ruth-nium (II), salts
    FE: And (Analytical reagent use); ANST (Analytical study); USES (Uses)
        (device for monitoring cells)
ΙΤ
    7631-86-9, Silica, analysis
    FE: AFU (Analytical role, unclassified); AMST (Analytical study)
    (device for monitoring cells)
59-05-2, Methatrexate | 151-21-3, Sodium didecyl sulfate, biblogical
TΤ
     studies 865-21-4, Vinblastine 7757-33-7, Scdium Sulfite /782-44-7,
    Omygen, biological studies 26628-22-8, Sedium Azide
                                                            35607-56-0,
     Cefoxitin : 69268-75-2, Cefuroxime : 55721-3:-1, Ciprofloxadi...
    EL: BSU (Biological study, unplassified); (IOL (Biological study)
        (device for monitoring cells)
    57-92-1, Streptomycin, biological studies 113-24-6, Sodium pyruvate
ΙT
     139:-39-3, Fungizone 1406-05-3, Penicillin 113978-18-6, Marrigel
     141:07-41-7, Matrix metalloproteinase
    FL: BUU (Biological use, unclassified); BLoL (Biological study; USES
     (Usus)
        THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
```

- (1) Bacon, J; Anal Chem 1997, V59(23), P2780 HCAPLUS
- (1) Berndt; US 6090574 A 2000
- (% Collins; US 6107083 A 2000
- (4) Gentlo; US 5906517 A 1999 HCAPLUS (5) Goswami, K; Fiber Optic Chemical Sensor for the Measurement of Partial Pressure of Oxygen 1988, V990, Pll1
- (m) Stitt; US 5567598 A 1906
- (7) Walt; US 5244636 A 1995 HCAPLUS
- (*) Wertz; US 444×634 A 1964
- (*) Wilfbeis, O; Mikrochimica Acta 1986, VN(5-6), 935 (HCAPLUS
- L79 ANSWER I OF L4 HUAPING COPYRIGHT 2002 ACS
- 2002:16:371 HCAPLUS $L\Pi$
- 110 136:196545
- Τī Method and apparatus for non-destructive screening of clinical specimen integrity
- T: Samscondar, James: Jacobs, Merrit Nylos
- One Telemetrik Iro., Mar., FVA.
- U.S., 23 pp., Cont.-in-part of U.S. Ser. No. 541,390, abandoned. S.() CODEN: USKKAM
- II. Patent
- Endlish L.-.
- T(ICM G01N033-48
- NCL 356040000
- CC 9-1 (Biochemical Methods)

FAN.CNT 1

PΙ

EATENT NO.	KIND	DATE	APELUCATION NO.	DATE
03 6753471		20020305	US 1997-871606	19970609 <

PEAL US 1098-941340 13951010 KH-В.

A method and app. for providing a non-destructive pre-test screen of specimen integrity for a blood analyzer by measurement of absorbance or reflectance is provided. The method involves measurement of polychromatic light in the near IE and adjacent visible region, which is either transmitted or reflected from a specimen as presented for measurement, and correlation of the measurement, on the basis of predetd.

algorithms, to the quantity of a known substance contained in the sample. The app. employs a spectrophotometer which emits radiation which is split into a beam which passes to a sample and a ref. beam, the beam returning from the sample and the ref. beam are variably combined and further sepa, into various components by means of a grating and focused ento a linear array detector. A microprocessor receives output from the array detector and performs calons. of conon.(s) of the known substance(s). The invention provides quality assurance for state-of-the art blood analyzers and automated labs. by pre-screening serum and plasma integrity, even where labels on the sample

container would normally interfere with a quality assurance assessment, identifying camples not suitable for sertain blood tests, or, if tests are conducted on specimens with compromised integrity, the pre-screening results will aid in the interpretation of the test results.

ST app screening clin specimen

ΙT Sensors

(Linear array: method and app. for non-destructive screening of clin. specimen integrity)

ΙT Light

(Folyanromatic; method and app. for non-destructive screening of clin. specimen integrity)

ΙT Analysis

> (elin.; method and app. for non-destructive screening of plin. specimen integrity)

ΙT Absorption spectroscopy

Algorithm

Analytical apparatus

ΙT

ΙT

ΙT

TΤ

ΤT

ΤT

Blood analysis Blood plasma Blood serum Concentration (condition) O ntainers Diffraction gratings Frequency Labels Light Mathematical methods Modecules Optical reflection Optical transmission Quality control Radiation. Samples Spectra Spectrometers Standard substances, analytical Time Turbidity TV and visible spectroscopy Wavelength (method and app. for non-destructive screening of clin. specimen integrity) Hemoglobins F.L.: ANT (Analyte); ANST (Analytical study) (method and app. for non-destructive screening of clin. specimen integrity) Computers (micriprocessors; method and app. for non-destructive screening of clin. specimen integrity) IR radiation IF. spectroscopy (near-IE; method and app. for non-destructive screening of clin. specimen integrity) Seybean cil RL: ANT (Analyte); ANST (Analytical study) (phospholip:a-stabilized; method and app. for non-destructive screening of clin. specimen integrity) 114-25-0, Biliverdin 635-65-4, Bilirubin, analysis RL: ANT (Analyte); ANST (Analytical study) (method and app. for non-destructive screening of clin. specimen integrity: RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD EΕ (1) Amen; CA 2019011 1994 (2) Congeshall; US 3702736 A 1972(3) Heinemann; US 8291884 A 1994 (4) Jacobs: US 5346492 A 1998 (b) Jacques; US 5313790 A 1994 (6) Karkar: US 5066659 A 1991 (T) Lundsquard; US 526:646 A 1334 (8) Lundsquard; US 5366903 A 1944 (9) McDeal; US 5734469 A 1998 (10) Potratz; US 5351685 A 1994 (11) Purdy; US 5360004 A 1994 LT9 ANSWER 3 OF 24 HEAPLUS COPYRIGHT 2002 ACS 2501:585529 HCAPLUS $E\Pi$ 1:6:17937 D11

System and method for analyzing antibiotic susceptibility of biological

```
samples
    Wiles, Timothy M.; Turner, David J.; O'connell,
    Michael A.; Parmigiani, Giovanni; Clyde, Merlise
PA
    Becton, Dickinson and Co., USA
    Eur. Pat. Appl., 32 pp.
ЗП
    CODEN: EPHKEW
DТ
    Fatent
    English
I.
     ICM G01N021-31
IC
0.1
    10-5 (Microbeal, Algal, and Fungal Biochemistry)
    Déstion pross-reference(s): 1
FAN.CUT 1
    FATENT NO.
                                         AFPLICATION NO. DATE
                    KIND LATE
    EP 1160564 A2 30011205 EF 2001-111418 20010510 <--
       E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, BO
                    A2 20020909
A 20000031 <--
                                          JE 2001-160395 20010529 <--
     TP 2002125697
PRAI US 2000-983891
   A system and method for analyzing samples, such as biol. samples, to
    accurately and effectively det. the susceptibility of the samples to
    antimicrobial materials, to det. min. inhibitory concn. (MIC)
    values for the resp. samples and antimicrobial materials. At each of a
    plurality of time intervals, the system and
    method directs a plurality of different analyzing light
    wavelengths, such as red, green and blue wavelengths,
    - nto each of a plurality of sample wells, and detects a resp.
    resultant light wavelength emanating from the resp.
    sample wells for each of the analyzing light wavelengths
     . The system and method uses resultant light
    wavelengths to generate at least two growth indicator
     characteristic curves representing, for example, the redox state
    and turbidity characteristics of the sample wells. The system
    then uses the redox state and turbidity
    -characteristics of sample wells contq. the same antimicrobial material to
    get, the MIC value for that material in relation to the sample contained
    on those wells.
    antibiotic susceptibility bidl sample
ST
\Gamma T
    Antibiotics
    Antimidrobial agents
      Computer application
      Drug screening
      Mathematical methods
      Measuring apparatus
      Redox potential
      Turbidity
       gsystem and method for analyzing antibiotic susceptibility of biol.
199 ANSWER : OF 04 HCAPLUS COPYRIGHT 2002 ACS
    _001:81/0:8 HCAPLUS
7
     139:528925
DD
    Method for non-invasive spectrophotometric blood oxygenation monitoring
T, I
\Pi\Pi
    Bernit, Paul
    Gas Medical Systems, Inc., USA
PA
30)
    FOT Int. Appl., 35 pp.
    CODEN: PIKKEZ
DT'
    Patent
LE
    English
     10M G01N
Τ :
    9-1 (Bi)chemical Methods)
FAN.CET 1
                                         APPLICATION NO. DATE
     PATENT NO. KIND DATE
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                             ____
    WG 2001034107
                      A2
                             20011104
                                            WO 2001-US13875 20010430 <--
PΙ
         W: AE, AG, AL, AM, AT, AU, AC, BA, BB, BG, BE, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DC, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KC, KE, KE, KE, LC, LK, LE, LS, LT,
             LY, LY, MA, MD, MG, ME, MN, MW, MX, M2, M0, M1, PL, PT, RO, RU,
             35, SE, SG, S1, SE, SL, TJ, TM, TE, PT, TZ, CA, UG, UZ, VN, YU,
             BA, ZW, AM, AB, BY, KG, KG, MG, EU, TJ, TM
         EW: AT, BE, CH, CY, DE, DH, EU, FI, FF, SB, GE, IE, IT, LU, MC, NL,
             PT, SE, TR, BE, EU, CE, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN,
             T19, T6
     AT 2001053255
                             2001111...
                                            AU 2001-59268
                                                            26010430 4--
                       A5
PRAI US 2000-201359P
                       5
                             2000051.
    Wo L001-U313875
                       1/
                            20010436
    A method and app. for non-invasively deta, the blood exygen sath, level
AB
    within a subject's tissue is provided that utilizes a near IR
     spectrophotometric (NIES) sensor capable of transmitting a light
     signal into the tissue of a subject and sensing the light signal
    ence it has passed through the tissue via transmittance or reflectance. The method includes the step of dotg, attenuation of the {\bf light}
     signal as the sum of: (1) attenuation attributable to decxyHb; (1i)
     attenuation attributable to dxyHk; and (iii) attenuation attributable to
     light scattering within the subject's tissue. The present method
     also makes it possible to account for attenuation attributable to fixed or
     ronst. light absorbing biol. tossue components, and attenuation
     attributable to variable characteristics of the sensor. By detg.
     differential attenuation as a function of wavelength, the
     attenuation attributable to tissue light scattering
     characteristics, fixed light absorbing components, and measuring
     app. characteristics are math. cancelled out or minimized
     relative to the attenuation attributable to decxyHb, and attenuation
     attributable to oxyHb.
ST
     moninvasive spectrophotometric blood exygenation remitoring
IT
     Information systems
        (data; method for non-invasive spectriphotometric blood oxygenation
        menitoring:
ΙΤ
     Afiral tissue
     Plood analysis
     Calibration
       Concentration (cond.tion)
       Light
       Light scattering
      Mathematical methods
     Optical absciption
       Optical reflection
       Optical transmission
     Oxydenation
       Wavelength
        (method for non-invasive spectrophotometric blood exygenation
        menitoring:
ΙΤ
     Remoglobins
     Hemoglobins, oxyhemoglobins
     FL: ANT (Analyte); ANUT (Analytical study)
        'method for non-invasive spectrophotometric blood oxygenation
        monitoring:
IT
     Sensors
        Thear IF spectrophotometric; method for non-invasive spectrophotometric
        blood oxygenation monitoring)
ΙT
     IR spectrometers
     IF spectroscopy
        (near-IR; method for non-invasive spectrophotometric blood oxygenation
        monitoring)
ΙT
     Gas sensors
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Foxygen; method for ner-invasive spectrophotometric blood oxygenation monutoring; 7781-44-7, Oxygen, analysis ΙT RL: ANT (Analyte); ANST (Analytical study) predisors; method for hom-invasive spectrophotometric blood oxygenation moniforing. L79 ANSWER 5 OF 24 HOAPLUS COPYRIGHT 2001 ACS 0001:649973 HCAPLUD AN 135:177(88 DMMethod and apparatus for determining the sensitivity of a microorganism to TI a growth altering agent Π I Wardlaw, Stephen C. EAUSA U.S., 9 pp., Cont.-in-part of U.S. 8,032,734. SO CODEN: USHKAM $D'I^{-}$ Paterit LAEnglish ICM 01.M001-16 IC 108 01.00001-18 NCL 435288700 9-1 (Bischemical Methods) Section pross-reference(s): 1, 10 FAN.CHT PATENT NO. KIND DATE APPLICATION NO. DATE US 1.016-477932 .0000105 k--P1US 60.1.7:4 A D0000068 PS 1999-256481 19990223 K-US 6140069 A D0001031 PS 1999-255681 19990223 K-NO L000004450 A D0001078 D0 L000-4480 .0000906 K-NO L001006048 A D0010708 D0 L001-43 .0010104 K-EF 1120467 A D0010708 BP L011-3000067 .0010108 K--]9990223 <--E: AT, BE, CH, DE, DK, ED, FR, GE, GE, IT, DI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO UF 2001218597 AU 20010614 CN 17/3886 A 20011128 ON 17/3986 A C0011128 PRAI US 1996-772169 P 19960307 US 1998-772175 P 19960307 US 1998-256681 AL 19990223 US 1998-256681 AL 19990223 W0 1999-724511 W 1 0990202 H--U3 1600-477932 A 10000108 H--A method and an app. for detg. the concn. at which a AВ growth-altering agent has an appreciable effect on the growth of a target microorganism are provided. The method Symposises the steps of (a) providing a microorganism growth medium; (h) providing a sensible reagent, which includes a growth -altering agent mixed with a marker that has a signal with a magnitude proportional to the concn. of the marker; (a) incorporating the sensible reagent into the growth medium, in a manner that creates a gradient of growth-altering agent and marker concns. within the growth meanum; (d) inoculating the growth medium with the target microorganism; (e) indubating the ing rulated growth medium for a period of time sufficient for the target mi moorganism to grow a detectable amt.; (f) -valuating growth characteristics of the microorganism in a region bonty, the growth-altering agent, (g) measuring the magnitude of the marker signal in that region; and (n) detg. the concn. of the growth-alterin; agent using the measured magnitude of the marker signal. app detg microorganism growth agent STΙΤ Molecules (Growth altering; method and app. for detg. sensitivity of a

microorganism to a growth altering agent) ΤT Fluorometers (Scanning; method and app. for detg. sensitivity of a microorganism to a growth altering agent) TΤ Antimicrobial agents Apparatus Concentration (condition) Culture media Growth, microbial Light scattering Mathematical methods Microorganism Mixing Sensors Time (method and app. for detg. sensitivity of a microorganism to a growth altering agent) TΤ Seapents EL: ANG (Analytical readent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (method and app. for detg. sensitivity of a microorganism to a growth altering agent; 20 THERE ARE 10 OFFED REFERENCES AVAILABLE FOR THIS RECORD P.E.CNT FΕ (1) Amon; EP 0635120 B1 1999 HCAPLUS (2) Buer; US 5547849 1996 HCAPLUS (+) B.ume; US 3925166 1975 (4) Ericsson; US 4778758 1988 (1) Ericsson; US FOLMERY 1991 HCAFLUS (r) Ericeson; TJ 56-9682 1997 () Kjellander: US 4204045 1980 (a) Lancaster; US 9501993 1996 HCAPLUS (m) McCoy: US 5700084 1997 ECAPLUS (10) Baskm; US 4790040 1986 (!1) Nishimura; US 5417959 1995 (12) Febertson; US 5206151 1995 ECAPLUS (13) Ochalkowsky; UP 4514495 1985 ECAPLUS (14) Schalkowsky; US 5346937 1993 (18) Schalkewsky; US 5863043 1996 HCAPLUS (16) Emith: US 4950455 1930 (17) Thempson; US 5164301 1992 HCAPLUS (18) Vesterberg; UN 4054400 1977 HCAFLUS (19) Warmlaw; DD 6/22734 2000 HCAFLUS i. 0) Wardiaw; US 6140069 2000 HCAPLUS ITE IMENTE S OF HA HOAPLUS COPYRICHT 2002 ACE L001:6496 \circlearrowleft HCAPLYS F.:135:177714 $I \in \mathbb{N}$ J. : Method for extending the range of an immunoassay 1:: Wei, The quan; Parkratz, Thomas John; Chu, Victor Pichai F_{T_1} Dade Behrung Inc., UDA W.S., 13 pp., Cont.-in-part of U.S. Ser. No. 166,026, abandoned. 50 CODEN: USMKAM UT Patent F.I English 10 DDM G01N033-53 TOS G01N033-543; G01N021-00; A61K049-00; C07K016-00 4350 (7100 NOL (. . . m-10 (Biochemical Mathada) FAN.CHT ! FATENT NO. KIND DATE APPLICATION NO. DATE _____ _____ ____ B1 20010904 US 1999-294489 19990420 <--US 6284472

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19981005 <--
PRAI US 1998-166006
                        B2
   - Calibrating an immunoassay by generating two reaction rate measuring
     curves, from samples having higher and lower relative levels of antigen,
     extrapolating a combination of the curves to cover sample concis
     . known to contain an excess of antigen relative to an amt. of capture
     reagent and sembining the low end linear potion of the higher reaction
     rate measuring curve with the higher end portion of the extrapolated
     reaction rate measuring curve, thereby eliminating measuring inaccuracies
     otherwise arising from the hook effect. For antigen concns.
     nigher than the assay range, a high antigen signal utilizing the two rates
     avoids reporting talms result:.
ST
     extending range imminoassay
     Froteins, specific or class
I T
     FL: ANT (Analyte,; ANST (Analytical study)
        (C-reactive; method for extending range of immunoassay)
] T
     Emagents
     FL: AFG (Amalytical reagent use); ANST (Analytical study); USES (Uses)
        (Capture; method for extending range of immunoassay)
     Calibration
       Concentration (deradition)
     Immuneassay
       Mathematical methods
     Reaction kinetics
       Regression analysis
     Samples
     Volume:
        (method for extending range of immunoassay)
     Antidens
ΤT
     RL: ANT (Analyte,; ANST (Analytical study)
        (method for extending range of immunoassay)
     Immunicassay
        (turbidimetric; method for extending range of immunoassay)
              THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
(1) Chadney; UC 55:4539 1096 HCAPLUS
(i) Cradle; US 4595661 1996 HCAPLUS
(A) Diamandis; US 5099425 1992 HCAPLUS
(4) Frengen; UN 5723346 1998 HCAPLUS
(to Frengen; UN 503364), 1998 HCAPLUS
(6) Graham; US 474584. 1868 HCAPLUD
(7) Hirai; US 5348368 1939 HCAPLUS
(8) Kappe: US 4055752 1977 HCAPLUS
(9) Kaspar; US 4960833 1930 HCAPLUS
(10) Lindme; UD 55-5241 1996
(11) Oh; UC 5583055 1996 HCAPLUS
(10) Ch; UC 57-335-1996 HCAPLUS
(13) Fouriquez; UC 41.3125 1979 HCAPLUS
(14) Somelli; UN 5 482830 1995 HCAPLUS
(15) Schafer; US 5420 42 1995 HCAPLUS
(16) Tung; US 4788138 1988 HCAPLUS
(17) Wu; US 4358852 1983
(18) Yamada; US 5253556 1995
LT9 ANSWER 7 OF L1 HCAPLUS COPYRIGHT 2002 ACS
I_{\bullet}!
     2001:995403 HCAPLUS
     135:149576
[di
Τ!
     Automated optical reader for multiple samples,
     especially for nucleic acid assays
     Andrews, Jeffrey P.; O'Eeefe, Christian V.; Scrivens, Brian G.; Pope,
TH
     Willard C.; Hansen, Timothy; Failing, Frank
Ρ...
     Becton, Dickinson and Company, USA
     Ear. Pat. Appl., 40 pp.
SU
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CODEN: EPXEDW

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DT
     Patent
    English
LA
\mathbb{I} \subset
     ICM G01N021-64
CC
     4-1 (Biochemical Methods)
     Rection cross-reference(s): 3
FAN.CHT 1
    PATENT NO.
                  KIND PATE
                                          APPLICATION NO. DATE
    EP 1124128 A2 20010816 EP 2000-1.3062 20001221
PΤ
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, SI, RO
→ TP 2001255272 A2 20010911
PRAI US 2000-483686 A 20000114
                                           JP 2001-6091 20010115
   An app. and method employ a plurality of light
    emitting devices which each can get light through a resp.
    optical fiber toward a resp. sample of a plurality of samples in
    a time-staggered manner. Light is generated in each
    of the samples at different times consistent with the
     times at which light is irradiated onto the sample. A
     single detector is used to detect the lights emitted from the
    plurality of samples at these different times. A
    plurality of bifurcated optical cable are coupled to the
     light emitting devices and single light detector, and
     the integrated end of each biturcated cable acts as the light
     emitting port and light detecting port. Multiple
     targets can be detected from each of the plurality of samples in
     the same manner by providing an app. and method employing a different
     plurality of light emitting devices and single detector
     for each tardet to be detected.
    automated optical reader app nucleic acid assay;
     multiple sample automated analysis app
    :amples
ΙT
        (anal. of multiple; automated optical reader for
        multiple samples, esp. for nucleic acid assays)
ΙΤ
     Analysis
     Process automation
        (automated anal.; automated optical reader for
        multiple samples, esp. for nucleic acid assays)
TT
     Algorithm
       Electroluminescent devices
     Fluius
       Light sources
       Optical cables
      Optical detectors
      Optical fibers
     Photomultipliers
        (automated optical reader for multiple samples,
        esp. for nucleic acid assays)
IΤ
    Nucleic acids
     KL: ANT (Analyte); ANST (Analytical study)
        (automated optical reader for multiple samples,
        esp. for nucleic acid assays)
     Probes (nucleic acid)
ΤT
     HL: AEG (Analytical reagent use); ANST (Analytical study); USES (Uses)
        (automated optical reader for multiple samples,
        esp. for nucleic acid assays)
TT
     Analytical apparatus
        (automated; automated optical reader for
        multiple samples, esp. for nucleic acid assays)
ΙT
     Computers
        (microcomputers; automated optical reader for
        multiple samples, esp. for nucleic acid assays)
     Microtiter plates
ΙΤ
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(microwell arrays; automated optical reader for multiple samples, esp. for nucleic acil assays

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L79 ANSWER 8 OF 24 HOAPLUS COPYRIGHT 2002 ABS
    2001:417342 HCAPLUS
MA
     135:16356
DH
    Method of measuring tissue nemoglobin saturation using gaussian
T.
    decomposition:
IM
    Wilson, David A.
FA
    Johns Hopkins University, USA
    PCT Int. Appl., 76 pp.
S(\cdot)
    CODEN: FINKED
\Gamma^{T}
    Fatent
LL
    English
     ICM G01N021-35
IC
     103 A61B005-00
    +-5 (Biochemical Methods)
CC
FAN.CMT 1
                                         APPLICATION NO. DATE
    FATENT NO.
                    KIND DATE
     F^{*}\mathbb{I}
    WO 2001040776
        W: AE, AG, AL, AM, AT, AC, AL, BA, BB, BG, BE, BY, BS, CA, CH, CN,
            CE, CU, CZ, DE, DK, DM, DC, BE, EJ, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JF, KE, KG, KP, KE, KC, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MH, NW, ME, ME, NO, MH, PL, PT, RO, RU,
             SD, GE, SG, SI, SK, CD, TI, TH, TR, TT, TD, UA, UG, UB, UZ, VN,
             YU, CA, EN, AM, AZ, BY, KG, KS, MD, KU, TJ, TM
         HW: GH, GM, KE, LS, MW, MC, SD, SD, SD, TE, FG, EW, AT, BE, CH, CY,
            DE, DE, ES, EI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
PU, CF, CG, CI, CM, GA, CU, CW, Mb, ME, ME, SU, TD, TG
PHAI US 1999-1005539 P 19991202 0--
    The constituents of perebual tissues that contribute to light
    absorbency, i.e., oxyHb, decwyHb, water, fipid, cytochrome oxidase and a
    component for characterizing light loss due to scattering, are
    further characterized and used to construct a model system that emulates
    cerebral tisque reflectance spectra in a mariety of conditions. Using
    this model system in a reverse mode, compd. spectra collected from brain
    tissue are decompo, into individual spectra features. The values for
    leatures attributable to oxyMb and deoxyMb are then used to construct a
    ratio that quantifies the percentage of total Hb that contains exygen.
    Recause the major portion of light, collected by the detecting
    element of the equipment has transited through brain dissue, this ratio
    Lecomes a quant, measure of brain tissue Hb sath. The decompn. anal.
    method is generally applicable to a variety of tissues besides brain
    tissue.
    tissue Hb sath gaussian decompn
ST
ΙT
     Ereray
        (Light; method of measuring tissue Hb sath. Using gaussian
        descents.)
IΤ
    Arimal tissue
    Apparatus
    Brain
    Hatabases
      Light
      Light scattering
      Mathematical methods
    optical absorption
    omygenation
     Reflection spectra
        (method of measuring tissue Hb sath, using gaussian decompn.)
IΤ
     Hemo rlobins
     Hemoglobins, exyhemoglobins
     EL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical
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study); BIOL (Biological study)
        (method of measuring tissue Hb satr. using gaussian decompos.)
     Lipids, biological studies
ΙΤ
     PL: ESU (Biclinginal study, unclassified); PFP (Properties); EICL
     Piological study.
        (method of measuring tissue Hb sath, using gaussian decompn.)
ΤT
     IF. spectroscopy
        (near-IE; method of measuring tissue Bb sath, using gaussian decompn.)
     7782-44-7, Oxygen, analysis
ΙT
     EL: ANT (Amalyte); ESU (Biological study, unclassified); PEP (Physical,

    nginesering in chemical process); ANST (Analytical study); BIOL

     (Elological study); PROC (Process)
        (method of measuring tissue Hb sath, using gaussian decompa.)
     7732-18-5, Water, biological studies | 9001-18-5, cytochrome oxidase
ΙT
     FI: ESU (Biological study, unclassified); PFP (Properties); BICL
     (Biological study)
        (method of measuring tissue Hb sath, using gaussian decomps.)
PE.ONT 6 THEFE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
(1) Bern:, P: PROCEETINGS OF THE ANNUAL NORTHEAST BIGENGINEERING CONFERENCE
    1935, VOONE 21, E105
(2) Gleve, E: TENTILVEREDLUNG 1995, V30(7/03), P109
(s) Lepper, J; US 5743162 A 1996
(4) Mannheimer, F; US 0782237 A 1933
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(G) Simeriman, K; APELIED SPECTROSCOPY 1999, V55(E), P:25 HCAPLUS
179 ANSWER FOR 24 HOAPLES COPYRIGHT 2012 ACS
    20001:376867 HCAPLTS
Z11
E:11
     134:75025.
    Method and scattered light-measuring apparatus for measuring a
ΤI
    scattered light and method of urinalysis using the same
III
    - Mawamura, Tatsuriu
    Matsushita Electric Inquatrial Co., Ltd., Japan
FΑ
SO
     Eur. Pat. Appl., 11 pp.
     CODEN: EPHHOW
E'T
    Fatent
LA
     English
    ICM G01N021-51
IC
Citi
     9-1 [Biochemical Methods]
     Section ordestreferonce(s): 78
FAN.CHT 1
     PATENT NO. KIND DATE
                                          APPLICATION NO. DATE
     EP 1102059 A1 20010818 EP 2000-125003 20001118 C
Er At, et, ct, tE, cm, Er, FF, GB, GA, CT, LI, LU, NL, SE, MC, PT,
ΡI
            IE, SI, LI, LV, FI, EO
     лр 1999-31:78с — АЛ 19991113 <--
                                           JP 2000-321074 20001020 K--
PHAI JF 1999-31:786
     The present invention provides a method and an app. which eliminate the
     influences of a smattered light arising due to the pollutants
     inside and on the surfaces of an optical window, differences in refractive
     index and light transmittance of a soln, to be detected, and the
     enstruction are to suspending particles and the like to achieve a
     measurement with high precision and high practicability in the measurement
     of the abattered light. The adattered light
     propagating within a prescribed angle perpendicularly to the direction of
     propagation of the light to be propagated through the inside of
     the solm. is measured. Further, the position of the optical axis of the
     light to be propagated through the inside of the soln, and/or the
     position of the photosensor in the direction of the optical axis are set
     so that the influence of the scattered light arising at and on-
     the purface of the optibal axis is not more than a predetd, value within a
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practically allowable range. The protein concn. in urine was
    detd. by measuring turbidity after heat treatment.
    scattered light measuring app urinalysis; protein urine
ST
    scattered light analysis
ΙT
    Proteins, general, analysis
    FL: ANT (Analyte); PEP (Physical, engineering or chemical process); ANST
     (Analytical study); FEOC (Process)
        (coagulation in urine for detg. protein concn.; method and
       scattered light-measuring app. for measuring scattered
       light and method of armualysis using same
    Fartisles
TΤ
        (interfering; method and stattered light-measuring app. for
       measuring scattered light and method of urinalysis using
       Same:
    UV and visible spectroscopy
ΙT
       (light-scattering; method and scattered light
        -measuring app. for measuring scattered light and method of
        urinalysis using same:
    Light scattering
      Mathematical methods
      Polarized light
      Refractive index
     Trine analysis
        (method and scattered light-measuring app. for measuring
        scattered light and method of urinalysis using same)
TΤ
    Feagents
    FL: AEG (Analytical readers use); ANST (Analytical study); USES [Uses]
        (method and scattered light-measuring app. for measuring
        scattered light and method of urinalysis using same)
ΤT
    Optical instruments
        (scatterometers; method and scattered light-measuring app.
        for measuring scattered light and method of urinalysis using
        same)
RE.CNT 4
             THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
F.E
(1) Barber, D; US E 440178 A 1 +4 : HCAPLUS
(1) Canon Ex; EF 0495125 A 1991 BCAPLUS
(3) de Maeyer Leo, C; US 4070420 A 1973
(4) Kowa Co; EP 0361770 A 1944
L79 ANSWER 10 OF 04 HCAPLUS COPYRIGHT 2002 ACS
    2000:807747 HCAPLUS
AI:
     188:331765
DI
    Device and procedure for the monitoring and control of microorganism
ΤI
    populations in biologically active fluids
    Höefrer, Fromas; Holzhauer, Peter; Walitza, Eckehard
II:
    Fraunhöter-Gesellschaft zur Foerderung der Angewandten Forschung EV,
FV.
    Germany
    Ger. often., 12 pp.
50
    CODEN: OWKEBE
ÐΤ
    Fatent
LÀ
    German
    10M - 01120001 - 02
TO
CH
    9-1 (Blockemical Methods)
FAN.CNT 1
    FATENT NO.
                 KIND DATE
                                          APPLICATION NO. DATE
    DE 19931939
                     A1 2.0001116
                                           DE 1999-19921999 19990512 <--
PΙ
    Wo 2000070073
                     A2:
                           20001123
                                           WO 2000-EP4289 20000512 <--
                    АЗ
     WO 2.00070073
                           2 (010301
        W: CA, JP, JS
         HW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
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PT, SE

A2 2002021 -EP 2000-936736 20000512 <--EP 1179174 P: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI 1999081. <--PRAI DE 1:49-19921999 A WO 2-00-EP4289 Į√J 2000012 < --A data acquisition mechanism and subsequent data processing are used to A.E. monitor metabolic parameters in a biol. active fluid so as to detn. the concn. of organisms. Thus, metabolic products or substrates such as carbon dickide, nydrogen, oxygen, etc., may be monitored to define bacterial or fungal populations in fluids. migreorganism monitoring metabolite liq control app STΙT Algorithm Pacillus (bacterium genus) Bacillus subtilis Candida Clostriaium Control apparatus Data processing Desulfatomaculum Electric conductivity Enterobacteriaceae Enterococcus Escherichia codi Laptobacillus Leadonestor Liquids Methanobacterium Methanococcus Midroccabus Eseudomanas Eseudomonas aenuginosa Redox potential Sadonarimydes Sardina Sensors Staphylococcus St rept occordus ĸН (mervice and procedure for maniforing and control of microorganism populations in bidl. active fluids) ΙT Computers (macroprocessors; device and procedure for monitoring and control of misroorganism populations in biol. active fluids) Aerobio bacteria ΙT (spore-forming; device and procedure for monitoring and control of microcipanism populations in biol. active fluids) 53-21-5, Lactic soid, analysis 64-17-5, Ethanol, analysis 64-18-6, Formir soid, analysis 64-18-7, Acetic soid, analysis 67-64-1, Acetone, TT analysis 71-23-8, Propanol, analysis 71-36-5, Butanol, analysis 74-82-8, Methane, analysis 107-92-6, Butyric acid, analysis 124-38-9, Carbon didxide, analysis 1838-74-9, Hydrogen, analysis 3812-32-6, Carbunate, analysis 7064-41-7, Ammonia, analysis 7727-37-9, Nitrogen, analysis 7762-44-7, Oxygen, analysis 7763-06-4, Hydrogen sulfide, analysis 14797-55-8, Nitrate, analysis 14797-65-0, Nitrite, analysis 14795-03-9, Ammonium, analysis 18496-25-8, Sulfide EL: ANT (Analyte); ANST (Analytical study) estervice and procedure for monitoring and control of microorganism populations in biol. active flaids) THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 4 F.F.

(1) Anon; DE 19605755 A1 HCAPLUS (2) Anon; DE 4415444 A1 HCAPLUS

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L79 ANSWER 11 OF 24 HCAPLUS COPYRIGHT 2002 ACS
     2000:790744 HCAPLUJ
AII
     133:419314
\Gamma
     A combined rapid anti-microbial susceptibility assay and microorganism
ΤΙ
     identification systems
I:1
     William, Greatry B.; Nothaft, Damiel; Enside, Glenn F.; Burther, Kathleen
     M.; Handas, M nte
PA
     Dade Microscan Inc., USA
     FOT Int. Appl., 50 pp.
SU
     CODEN: PIMMEC
DT
     Patenit
ĽΑ
     English
TC
     G01N035-02; C12M001-34; C12M001-20
     9-16 (Biodhemadai Methods)
\mathbb{C}\mathbb{C}
     Section arcss-reference(s): 1, 10
FAN.CHT 1
                                             APPLICATION NO. DATE
     PATERT NO.
                      HIND JATE
                                             ______
     ______
                                            WO 2016-0312781 20000501 :--
     W6 2000067037 AL 20001109 W6 2000067037 AS 20011011
ΡŢ
         W: AE, AL, AM, AT, AU, AS, BA, BE, BG, BE, BY, CA, CH, CN, CE, CU,
             CO, DE, DK, DM, EE, EO, FI, GE, GD, GE, GH, GM, HR, HU, ID, IL,
             IN, IG, JF, KE, RG, KP, ER, ED, LJ, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MM, DO, MC, PL, PT, RG, RU, SE, SE, JG, SI, SR, GL, TJ, TM, TR, TT, TG, WA, US, UZ, VN, YU, ZA, ZW, AN, AZ,
             BY, KG, KD, MD, RU, TJ, TM
         HW: GH, GM, KE, LS, MW, SD, CH, CC, TÜ, CG, CW, AT, BE, CH, CY, DE,
             DE, FC, FI, FR, GB, GE, IE, IT, LT, MC, ML, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, CN, GW, ML, ME, NE, SN, TD, TG
                                                               20000501 ---
                      Ar 20020206 EP 2000-930840
     EF 1177448
         H: AT, PH, CH, DE, DK, ED, FH, GH, GH, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, EC
PFAI US 1499-181800P P 109904L9 ---
US 1999-177819P P 19990817 ---
     US 1000-595213 A 200004.0 > --
WO 1000-U012751 W 100005 1 > --
    In response to the need for highly-sensitive antibiotic susceptibility
AΞ
     assays and identification assays that do not require extensive incubation
     times, the present invention provides automated assay
     methods and systems that permit the detn. of antibiotic susceptibilities
     and/or microsigamism identification in a time frame that is
     substantially shorter than has previously been antainable using a hybrid
     system that combines turbidimetric and fluorescence
     detral using a single, Mear-plastic assay platform. Related devices,
     kits, and components thereof are also displosed.
ST
    microbial susceptibility assay microsiganism system
ΙΤ
     Colorimetry
        (Bichromatic; a combined rapid ant:-microbial susceptibility assay and
        microorganism identification system)
IΤ
     Computers
        (Central processing units; a combined rapid anti-microbial
        susceptibility assay and microorganism identification system)
IT
     Optics
        (Multiple wavelength; a combined rapid
        anti-micropial susceptibility assay and microproganism identification
        system)
ΙT
        (Multiwell; a simplified rapid anti-microbial susceptibility
        assay and microorganism identification system;
ΙΤ
     Plates
        (Plastic sample; a combined rapid anti-microbial susceptibility assay
```

and microorganism identification system)

IT Algorithm

Analytical apparatus

Antibiotics

Antimicrobial agents

Apparatus

Color

Colorimeters

Colorimetry

Computer application

Culture media

lives

Enterobacteriaceae

Fluorescent substances

Fluorometers

Fluorimetry

Gram-negative bacteria

Gram-rositive bacteria (Firmicutes)

Inks

Interface

Liquids

Microorganism

Mixina

Fairts

Suspensions

Temperature

Test kits

Time

Turbidimetry

(a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Feagents

FL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Flastics, uses

RL: NUU (Other use, unclassified); USES (Uses)

(a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Analysis

Process automation

(automated anal.; a combined rapid anti-microbial

suspeptibility assay and microorganism identification system)

IT Construction materials

(recards; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Light

(fluorescent; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Wells

(multi-; a combined rapid anti-microbial susceptibility assay
and microorganism identification system)

IT Opacity

(opacification; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Laboratory ware

(reaction vessels; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Containers

(reaction; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT Hydration, chemical

(rehydration; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

ΙT 9035-73-8, Oxidase

FL: AFG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BICL (Biol.gical study); USES (Uses)

(a combined rapid anti-microbial susceptibility assay and microorganism identification system)

- L79 ANSWER 12 OF 04 HCAPLUS COPYRIGHT 2000 ACS
- 2000:697799 HCAPLUS A:I
- 134:41258 [:]
- Τ... Evaluation of turbidity: correlation between Kerstez turbidimeter and nephelometric turbidimeter
 - Collado-Pernandez, M.; Gonzalez-Samjose, M. L.; Pino-Navarro, R.
- Department of Biotechnology and Food Schence, University of Burgos, (0,1)Purgos, Spain
- Food Chemistry (2000), 71(4), %63-160 50 CODEN: FOCHOJ; ISSN: 0303--146
- Eib Elsevier Schende Ltd.
- Journal E'T

ДH

- Λ . Enclish
- CC17-1 (Food and Feed Chemistry)
- Turbidity is a quality parameter that has an important role in Lfield lig. acceptance. Cloudiness of beverages and covering lig. are a consequence of manuf, processes and storage conditions. Spanish legislation defines the covering lig. turbidity in canning by Kerstez turbidimeter units (KTU), which is a sensorial measure. It is necessary to find a correlation between sensorial and instrumental measurements. This work studied the relationship between KTU and normalometric turbidimeter units (ETU) and established a math. model, which allowed the expression of the turbidity of lig. products in KTU frim measurements in nephelometric turbidimeter units. This math. model corresponds to a non-linear simple correlation model (RTU-NTU). The best adjustment was a Reciprocal-Y model.
- ζ: " food analysis turbidimetry reprelemetry
- 1: Food analysis

Nephelometry

Simulation and Mudeling, physicochemica.

Turbidimetry

(correlation between Kerstez turbidimeter and nephelometric turbidimeter in food anal. I

17 Measuring apparatus

Optical instruments

(turbidimeters; derrelation between Manstez turbidimeter and nephel metric turbidimeter in food anal.)

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD F.E. CNT 15 F.E

- (1) Andin, C; Food Chemistry 1996, V58(1), PJ41 HCAPLUS
- (L) BOE; Normas de datidad de okinservas vegetales 1984, 287, 288 and 289, 30-KI, 1-XII and 2-KII
- (s) Calvo, C; Información tecnica general 1971, 55, Pl
- (4) Calvo, C; Pevista de Agroquímica y Tecnología de Alimentos 1980, V20(1), 2144
- (b) Dickinson, E: An introduction to fold colloids 1992
- (6) Dickinson, E; Food Chemistry 1994, US1, PB4S HCAPLUS
 (7) Daran, L; Fevista de Agroquemica y Pechologia de Alimentos 1976, V16(1), Pa∃
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- (3) Genovese, I; Journal of Foot Science 1997, V62(6), P1171 HCAPLUS
- :10) Hernandez, E: Journal of Food Science 1931, V56, P747
- [11] Kramer, A; Food Technology 1969, V23, P92

- (12) Markowski, J; Fruit Processing 1998, V7, P277
- (13) Martin Belloso, O; Temas de tecnologia de conservas vegetales 1990, P87
- (14) Martine: Baigorri, E; Conservas vegetales 1384, 5, P9
- (15) Primo Yufera, E; Quimiba agrabbla 1987, V3, P878
- L79 ANSWER 15 OF 24 HCAPLUS COPYRIGHT 2002 ACS
- AN 2000:638185 HCAPLUS
- DN 133:218433
- TI Computerized method and apparatus for analyzing nucleic acid assay reading
- IN Yang, Harry, Schwarz, Daniel L.; Empres, Christopher M.; Moore, Richard L.; Harland, Perry D.; Johnson, Paula V.
- PA Becton, Dickinson and Company, USA
- SO Jpn. Koka: Tokkyo Koho, 64 pp. CODEN: JREKAF
- DT Fatent
- LA Japanese
- IC | CM | G01N033-50

108 | C120001-68; C12NC19-09

CC 3-1 (Biochemical Genetics)

FAN.CHT 1

	PATENT NO.	KIMD	DATE	APPLICATION NO.	DATE		
ΡI	JP 2900249701	F_{Δ} .	20000914	JP 1999-329984	19991119		
	US 6.36949	81	20010417	US 1998-196123	19981120		
PRAI	US 1993-1961 <u>-</u> 3	F_{λ}	19981120				

A computerized method and app. are disclosed for analyzing numerical data pertaining to a sample assay comprising at least one biol. or chem, sample. The data include a set of data pertaining to each resp. sample, with each set of data including a plurality of values wach representing a condition of the sample at a given time. The method and app. assign a resp. numerical value to each of the data values, math. combine the numerical values to deserate a total value, compare the total value to a threshold value, and control the system to indicate whether the sample has a predetd, characteristic based on a result of the comparison. Erior to calcn. of the sample value, filtering, normalizing and other correcting operations can be performed on the data to correct anomalous values in the data which could adversely affect the accuracy of the results. The method and app. perform the described functions by representing the data values as points on a graph having a vertical axis representing the magnitudes of the values and a horizontal axis representing a period of time during which readings of the sample were taken to optain the data values, identifying points on the graph having an anomalous characteristic, and correcting the anomalous points to produce a conceded plot of points on the graph, with each of the points of the cor. plot representing a magnitude of a corresponding one of the values. An area value is then calcd. which represents an approx. area between at least a portion of the cor. plot of points on the graph and the horizontal axis. The area value is compared to a threshold value to det, whether a certain condition exists in the sample to which the set of data pertains. Diagrams describing the app, assembly and the operation flow are given.

ST computer analyzer nucleic acid assay reading

IT Analysis

Analytical apparatus

(blockhem.; computerized method and app. for analyzing nucleic acid assay reading)

IT Computer application.

Jambies

(computerized method and app. for analyzing nucleic acid assay reading)

IT ::upleid abids

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical

```
study); BIOL (Biclogical study)
        (computerized method and app. for analyzing nucleic acid
        assay reading)
ΙΤ
    Information systems
        edata; computerized method and app. for analyzing nucleic
       arid assay reading)
L79 ANSWER 14 OF 24 HCAPLUS COPYFLIGHT 2002 ACS
    :000:313432 HCAPLUS
ΑN
     133:114329
EM
ΤΙ
    Multivariate statistics for energy-dispersive x-ray
    fluorescence analysis of chemical substances
    Henrich, Alexander: Itzel, Hans-Helmut: Hoffmann, Peter: Ortner, Hugo
IM
FA
   Merck Patent G.m.E.H., Germany
    FOT Int. Appl., 32 pp.
SO
    CODEN: PIKKOD
E-C
    Patent
LÀ
   German
TO
    ICM G01N023-20
CC
    79-2 (Inorganic Analytical Chemistry)
    Section cross-reference(s): 74
FAN.CHT 1
                                        APPLICATION NO. DATE
    PATENT NO. HIND DATE
    ______
                                         -----
    WO 2000043761 A2 20000717
                                        WO 2003-EP70 20030107 <--
    Wo 2000043761
                    A3 20001150
        W: JP, US
        EW: AT, PE, CH, CY, DE, DK, EC, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            ET, SE
                                    DE 1999-1992101, 1....
ER 2000-901071 20000107 K--
TITUMIN SE. MC,
                                         DE 1999-19921317 19990508 <--
                     DE 14921:17
                     EP 11449a6
        H: AT, BE, CH, DE, DK, EN, FH, GB, GH, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
                        13990123 ---
FFAI DE 1999-19901617 A
    DE 1499-19931317 A 19990508 --- WO 1000-EP70 W .3000107 ---
   A method was described for classifying and identifying, using
AΒ
    energy-dispersive x-ray fluorescence anal., onem. substances
    that have x-ray fluorescence lines that cannot be detected and
    which therefore cannot be classified by energy-dispersive x-ray
    fluorescence anal. alone. The method is characterized in that the
    sample to be analyzed is analyzed in its original packaging or natural
    state without prior processing in a sample vessel. Using this method, the
    sample is: (1) positioned in front of the measuring aperture in a sample
    chamber of an x-ray fluorescence app., (?) measured, and (3)
    classified and identified by application of multivariate,
    statistical techniques to the measurement signals obtained (i.e., to the
    Compton and Fayleigh scattering).
   energy dispersive x ray fluorescence; Compton scattering energy
ST
    dispersive x ray fluorescence; Rayleigh scattering energy
    dispersive x ray fluorescence; multivariate statistics
    energy dispersive x ray fluorescence
ΙΤ
    Light scattering
        (Rayleigh; multivariate statistics for energy-dispersive
        x-ray fluorescence anal. of chem. substances)
    X-ray fluorescence spectrometers
        (energy-dispersive; multivariate statistics for
        energy-dispersive x-ray fluorescence anal. of chem.
       substances)
    Meray spectroscopy
ΙΤ
    M-ray spectroscopy
        (fluorescence, energy-dispersive; multivariate
        statistics for energy-dispersive x-ray fluorescence anal. of
```

```
chem. substances)
ΙT
    Gompton effect
       Multivariate analysis
        (multivariate statistics for energy-dispersive x-ray
        fluorescence anal. of cnem. substances)
ΙT
     Fluorometry
     fluorometry
        (x-ray, energy-dispersive; multivariate statistics for
        energy-dispersive x-ray fluorescence anal. of chem.
        Eubstan mes)
     14:-35-9, Jodium cyanide 151-60-3, Potassium cyanide 471-34-1, Calcium
ΙT
     parmonate, properties 497-19-6, Sodium carbonate, properties 506-87-6,
     Ammonium carbonate 584-08-7, Potassium carbonate 1308-28-9,
     Chromium oxide (Cr203), properties 1309-37-1, Ferric cxide, properties 7439-40-6, Iron, properties 7447-40-7, Potassium chloride,
     properties 7647-14-5, Sodium chloride, properties 7661-49-4, Sodium
     fluoride, properties 7757-62-6, Sodium sulfate, properties 7778-50-9, fctassium dichromate 7778-5.-5, Potassium sulfate, properties
     7781-63-0, Ferrous sulfate heptahydrate 7783-10-2, Ammonium sulfate,
     rroperties 7783-85-9, Sulfurio acid, ammonium iron(2+) salt (2:2:1), hexahydrate 7787-96-6, Sulfurio acid, beryllium salt (1:1), tetrahydrate
     7783-98-9, Ammonium chromate ((MH4)2CrO4) 7788-99-0, Sulfurid
     acid, chromium(3+) potassium salt (2:1:1), dodecanydrate 7753-23-3, Fotassium fluoride 7791-18-6, Magnesium chloride, hexahydrate
     10025-77-1, Ferric chloride hemahydrate | 10034-99-8, Magnesium sulfate
     heptahydrate 10036-04-8, Calcium chloride dinydrate 10043-35-3, Boric
     acid (H?BO)), properties 10000-12-5, Chromic obloride
     hemahydrate 10101-41-4, Calcium sulfate dibydrate 12125-00-9, Ammonium
     chloride, properties 13943-58-3, Potassium ferrodyanide 14459-95-1,
     Potassium ferrodyanide trihydrate
     RL: FRP (Properties,
        (test substance; multivariate statistics for
        energy-dispersive x-ray fluorescence anal. of chem.
        substances)
L79 ANOWER 15 OF 24 HOAFLOO COETRIGHT 2002 ACG
AN
    -2000:161536 HCAFLUD
DN
    131:131:398
    Applicative and method for readentless analysis of biological samples
TI
1:1
    Jeng, Tzyy-wen; Mo, Dowell Larry L.; Pezzaniti, Joseph L.; Oosta, Gary M.;
    Shain, Eric P.
FΑ
    Abbott Laboratories, USA
SO
   FCT Int. Appl., 103 pp.
    CODEN: BIKKDI
DT
    Fater.t
LA.
    English
    ICM G01N021-27
IC
     ICS G01N021-41; G01N021-05
CC
     9-1 (Riochemical Methods,
FAN.CNT 1
     FATENT NO. KIND DATE
                                           APPLICATION NO. DATE
     -----
    WO 1999-UR19532 19990327 KH-
PΙ
                      As 20010222
     WO 2000013002
         W: CA, JF
EW: AT, BE, CH, CY, DE, DM, ES, FI, FE, GB, GE, IE, IT, LU, MC, NL,
             PT, CE
                            .00000711
     US +987182
                                            US (1996-141463 19960927 K--
                       EP 1999-942505 19990827 4--
     EP 1110075
         H: AT, BE, CH, DE, DK, HJ, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, FI
                                           US 1999-407397 19990928 <--
                       B1 200204 C
     US 6565109
                     A 19950627 <--
PRAI US 1398-141463
```

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19990927 <--
     WO 1939-US19532
                      W
     App. and method are disclosed for detg. at least one parameter, e.g.,
AΒ
     concn., of at least one analyte, e.g., area, of a biol. sample,
     e.g., trine. A biol. sample particularly suitable for the app. and method
    of this invention is urine. In general, spectroscopic measurements can be used to quantify the concns. of one or more analytes in a biol.
     sample. In order to obtain concn. values of certain analytes,
     such as Hb and bilirubin, visible light absorption spectroscopy
     can be used. In order to obtain concn. values of other
     amalytes, such as urea, preatinine, glubose, ketones, and protein, IR
     light absorption spectroscopy can be used. The app. and method of
     this invention utilize one or more math, techniques to improve
     the auguracy of measurement of parameters of analytes in a biol. sample.
     The invention also provides an app. and method for measuring the
     refractive index of a sample of biol. fluid while making spectroscopic
    measurements substantially simultaneously.
    app reagentless analysis biol sample; spectrometry biol fluid reagentless
ST
    analysis; refractive index analysis app biol fluid
ΤТ
    Absorption spectroscopy
    Biological materials
    Bloom analysis
     Eddy fluid
    Cerebrospinal fluid
    Electric impedance
    Fluoremetry
    IR spectroscopy
     Tens
      Light scattering
      Mathematical methods
     Faman spectroscopy
      Refractive index
     Saliva
    Spectioscopy
    Spratrani
     Sweat
     Temperature
    Urine analysis
        (app. and method for reagentless anal. of biol. samples)
TΤ
    Albumins, analysis
     Mitrites
     FL: ANT (Analyte); ANST (Analytical study)
        (app. and method for reagentless anal. of biol. samples)
工工
    Hemoglobins
     Retones, analysis
     Protoins, general, analysis
     EL: ANT (Analytical study); BIOL
     (Biclogical study); USES (Uses)
        (app. and method for reagentless anal. of biol. samples)
TΤ
     Spectrometers
        (colls, sample cell assembly; app. and method for reagentless anal. of
        bi:1. samples)
ΙT
     Readents
     FL: AFG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (container for, for materials not having significant spectral
        signature; app. and method for reagentless anal. of biol. samples)
ΙT
     Spectroscopy
        (deriv.; app. and method for reagentless anal. of biol. Samples)
ΙT
        (dialyzate of; app. and method for reagentless anal. of biol. samples)
ΙT
     Photometry
        (filter; app. and method for reagentless anal. of biol. samples)
```

```
ΙΤ
     Analytical apparatus
         (for measuring refractive index; app. and method for reagentless anal.
         of biol. samples:
ΙΤ
     Containers
         ,for reagent(s) for materials not having significant spectral
         signature; app. and method for reagentless anal. of biol. samples)
     Body fluid
ΙT
        sinterstitual; app. and method for reagentless anal. of biol. samples)
     UV and visible spectroscopy
IT
         (light-scattering; app. and method for reagentless anal. of
         Equal. samples)
ΙT
     Moise
        (method for redn. of; app. and method for reagentless anal. of biol.
         samples)
     50-99-7, Glucose, analysis 57-13-6, Urea, analysis 61-27-5, Creatinine
ΙΤ
     635-65-4, Bilarubin, analysis
     FL: ANT (Analyte); THU (Therapeutic use); ANSI (Analytical study); BIOL
      (Biological study); USES (Uses)
         (app. and method for reagentless anal. of biol. samples)
ΙΤ
     7732-18-5, Water, miscellaneous
     FL: MSC (Miscellaneous)
         (subtraction of absorption spectrum for; app. and method for
         reagentless anal. of biol. samples)
L79 ANSWER 16 OF 14 HOAFLUS COPYRIGHT 2002 ACS
     0000:9313L HCAPLUS
11A
[i]1
     132:131430
     Nonlinear optical scattering with imaging and fractal analysis for
TΙ
     determining the concentration of a material in a scattering
     Jungmann, Holger: Schietzel, Michael
IN
    MBE G.m.b.H., Germany
Ger. Offen., 10 pp.
F_{K}
50
     CODEN: GWEKEK
DT
     Patent
LA
    Carman
     TCM G01N021-47
TO
     TCS G01N021-55; G01N021-17; G01J003-42
Ci
     79-2 (Inordanic Analytical Chemistry)
FAN.CHT 1
     FATENT NO. KIND DATE
                                               APPLICATION NO. DATE
     ______
                                                _____

    DE 19831424
    Al
    20000293

    DE 19831424
    C2
    20001228

                                                DE 1998-19831424 19930714 <--
F'1
     A spectroscopic propedure for detq. the concn. of a material
AB
     within , positive ing medium, consists of the following steps: (1) illumination of the medium with {\bf light} at a continuous
     wavelength, (.) measuring the emitted light at a certain
     direction of the medium, (3) letg. the emission of the emitted
     light as a function of the wavelength compared with a
     sid., (4) introducing an absorption-free known scattering medium into the
     -ptical path, (5) measuring the light emitted at the certain
     direction of the sample and the scattering medium, (6) detg. the emission
     of the light emitted from the sample and the scattering medium compared with the std., (7) imaging the emissions detd. Without and with the scattering mediums, (8) detg. the fractal dimension of the images, and (3) detg. the concn. of the substance from the fractal
     Himension. In this way, previous knowledge of the optical and quant.
     properties of the scattering medium is not necessary.
ST
     scattering spectroscopy fractal imaging gas sensor
ΤT
     Fractals
         (fractal dimension; nonlinear optical scattering with imaging and
         fractal anal. for detg. the concn. of a material in a
```

scattering medium)

IT Nonlinear optical properties

Nonlinear optical properties

(light scattering; nonlinear optical scattering with imaging and fractal anal. for detg. the concn. of a material in a scattering medium)

IT Gas sensors

Imaging

Light scattering

(nonlinear optical scattering with imaging and fractal anal. for detg. the **concn.** of a material in a scattering medium)

IT Light scattering

Light scattering

(monlinear; nonlinear optical scattering with imaging and fractal anal. for detg. the concn. of a material in a scattering medium)

RELONT 2 THERE ARE 2 CITED REPERENCES AVAILABLE FOR THIS RECORD

(1) Anon; EP 0810429 A1

(2) Anon: US 5588427

L79 ANSWER 17 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1998: 018864 HCAELUS

DN 129:227807

TI Method and apparatus for measurement of blood substitutes

IN Samscondar, James

FA Cme Telemetrix Inc., Can.

SO FOT Int. App.., 41 pp. CODEN: PIKKD2

DT Faterit

LA English

IC I M G01N021-27

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 6, 13

FAN.CNT 1

17111.				KIND	DATE	AFFLIC	ATION NO	D. D	ATE			
							· 					
F' 1	WO			Al KR, US	19980911	WO 199	97-CA753	1	9971016	<		
	FE	EW: A	Γ, BE,	CH, DE,	, OM, ES,						PT,	SE
	11	F: A	Γ, BE,		, OK, ES,						PT,	
	~ [E, FI 9242	Tr. Tr	20010904	JE 199	48-5390:"5	7 1	9971016	<		
					19970303	01 111			13/1013			
	W^{-1}	1997-03	475.4	W	19971616							

- Ab A method is disclosed whereby the concn. of a rhood substitute, such as cross-linked Hb, in a serum or plasma specimen is rapidly and accurately identified and quantified. The method includes making a spectrophotometric measurement of the blood substitute and calcg. the concn. based on a calibration algorithm. The method further takes the measured concn. of the blood substitute and uses it to correct for its effect, if any, on a measured analyte concn., e.g., serum/plasma total protein. Surther, the method arlows for the detr. of the concn. of true Hb in the presence of blood substitutes. The method is carried out in respect of samples contained in a primary or secondary labeled tube, or a pipet tip used to dispense serum or plasma in a blood analyzer.
- ST blook substitute analysis spectrophotometry

IT Prot-ins, general, analysis
EL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study,
unclassified); ANST (Analytical study); BIOL (Biological study); OCCU
(Occurrence)

(blood; method and app. for measurement of blood substitutes)

```
ΙT
     Hemoglobins
     PL: ANT (Analyte); BSU (Biblogical study, unclassified); THU (Therapeutic
     use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
        (crosslinked; method and app. for measurement of blood substitutes)
    Algorithm
     Flood
     Flood analysis
     Elect plasma
     Flood serum
     Eloud substitutes
     Hemolysis
     Fefluction spectrostopy
       Spectrophotometry
       Turbidity
        (method and app. for measurement of blood substitutes)
ΙT
    Pile pigments
     Frcteins, general, analysis
     Fb: ANT (Analyte); AEC (Analytical role, unclassified); BOC (Biological
     cocurrence); BSU (Biological study, unclassified); AMST (Analytical
     study); BIOL (Biological study); 0000 (Occurrence)
        (method and app. for measurement of blood substitutes)
ΤT
    Hemoglobins
     FL: ANT (Analyte); BOC (Biblogical becurrence); BSU (Biological study,
     unclassified); ANST (Analytical study); BIOL (Biological study); OCCU
     (Occurrence)
        (method and app. for measurement of blood substitutes)
     9000-97-9, Aspartate aminotransferase 9001-18-4, Creatine kunase
ΙT
     8001-00-9, Labtate dehydrogenase 9001-78-9, Alkaline phosphatase
     9046-.7-9, .gamma.-Glutamyltransferase
     FL: ANT (Analyte); ARU (Analytical role, unclassified); BAC (Biological
     activity or effector, except adverse); BOC (Biological occurrence); BPR
     (Biological process); BSU (Biological study, anclassified); ANST
     (Analytical study); BIOL (Billogical study); OCCU (Occurrence:; PROC
     (Process)
        (method and app. for measurement of blood substitutes)
     57-13-6, Orea, amalysis - 60-17-5, Oreatinine - 71-52-3, Bidarbonate,
ΙΤ
     analysis 114-25-0, Biliverdin 658-66-4, Bilirubin, analysis
     743 -96-4, Magnesium, analysis 7441-03-7, Potassium, analysis
     744 (-13-8, Sodium, analysis 7441-71-2, Calcium, analysis 16887-30-6,
     Chloride, analysis
     Fh: ANT (Analyte); ARU (Analytical role, unclassified); BOC (Biological
     occurrence;; BSU (Biological study, unclassified); ANST (Analytical
     study); BIOL (Biclogical study); OCCU (Occurrence)
        (method and app. for measurement of blood substitutes)
    197462-97-8, Hemolink
ΙT
     Fh: ANT (Ahalyte); BUU (Biological use, unclassified ; THU (Therapeutic
     use; AMST (Amalytical study); BIGL (Biclogical study); USES (Uses)
        imethod and app. for measurement of blood substitutes)
L79 AMSWER 18 OF ..4 HCAPLUS COFFRIGHT 2002 ACS
    1998:21756.: RCAFLUS
IIA
D11
    138:354:13
    Reaction time window detn. for rate assays using
TΙ
     turbidimetry and nephelometry
I11
    Patzke, Juergen
PΑ
    Fehringwerke A.-G., Germany
SO
    Ger. Offen., 18 pp.
    CODEN: GWXXBX
DT
    Patent
LA
    German
    ICM G01N037-00
ICS G01N033-557; G01N021-49; G01N021-75;
IC
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G01N021-82

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3-5 (Biochemical Methods)
     Jestion pross-reference(s:: 14, 15
FAN.CHT 1
                    KIND DATE
                                        APPLICATION NO. DATE
    PATENT NO.
     ______
                                          ______
                    Al
    DE 13040131
                           19920403
                                        DE 1996-19640121 19960928 <--
EP 1997-115225 19970903 :--
PΙ
                    A2 199804:1
    EF 833153
        R: AT, BE, CH, DE, DH, ES, FR, GB, CR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
                           . 0000323
                                         US 1997-956544
     11.1 6044330
                                                          19970924 -:--
                      A
    CA 2216495
                     AA 19980313
                                         CA 1997-2216895 19970926 :--
    JF 10111248
                     Ad 19480413
                                          ap 1997-277950 19970926 -:--
    0.1 6317702
                                          ÚS 2000-503152 20000211 <--
                     31
                           _1011113
PRAI DE 1990-19640121 A
                          1996ტმექ <--
    US 1997-936544 Al 19970924 <--
    The invention concerns a method to det. The time window for
AΒ
    measuring quantities that change rate during the progress of the reaction.
    The max. quantity Lat) is detu. in the linear region; values detd. in a
    tarst expt. are used in a second detn. to derive the time
    window. Also polynoms that fit the L(t) function can be used to
     calc. the time window. The method is used in
     turbidimetry, nephelometry and light scattering
    measurements for antibody-antique reactions, plasma proteins and blood
    edotting.
    reaction time window turbidimetry nephelometry;
ST
    clotting immuneassay protein time window
ΤТ
    Immuneqlobulins
    FI: ANT (Amalyte); ANST (Amalytical study)
        (A; reaction time window deth. for rate assays using
        turbidimetry and Lephe.ometry)
ΙT
     Fibrinoder dedradation products
     FL: ANT (Analyte); ANST (Analytical study)
        (DE: reaction time window detn. for rate assays using
        turbidimetry and nephelometry)
ΤT
     Froteins, general, analysis
     FE: ANT (Amalyte); ANST (Amalytical study)
        (blood; reaction time window detn. for rate assays using
        turbidimetry and nephelometry)
TT'
     Acialysis
       (clin.; reaction time window detn. for rate assays using
        turbidimetry and nephelometry)
ΤT
    Algorithm
    Blood coadulation
     Immunicassay
     Latex
    Nephelometry
       Turbidimetry
       (reaction time window beth. for rate assays using
       turbidimetry and nephelometry)
ΙT
    Ferriting
     Prostate-specific antique
     EL: ANT (Analyte); ANST (Analytical study)
        crearting time window lett. for rate issays using
        turbidimetry and neptelometry)
LT9 ANSWER 19 OF .4 HCAPLUL COPYRIGHT 2002 ACS
     1997:445079 HCAPLUS
ΑN
     1.7:63-58
DN
    E-agent system and method for the differentiation and identification of
ΤI
     reticalocytes
    Studnolme, Robert M.; Marchall, Paul N.; Embleton, Anne M.; Gläzier, John
ΙN
    G.; Van Howe, Luc
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Abbott Laboratories, USA

PΑ

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FOT Int. Appl., 28 pp.
    CHEN: PIMMED
    Eatent
LA
    English
     I:M G01N033-80
10
     9-4 (Biochemical Methods)
CC
     Section cross-reference(s): 13
FAN.CHT 1
                                          APPLICATION NO. DATE
    EATENT NO. KIND DATE APPLICATION NO. DATE

WO 0719:56 AT 10070509 WO 10:6-US184/1 10061118 <--
    EATENT NO.
                     KIND DATE
ΞI
        W: CA, JP
        PW: AT, BE, CH, DE, DK, E3, FI, FR, GB, GR, TE, IT, LU, MC, NL, PT, SE
    US 5753784 A 19980331 US 1995-500601 19951120 <--
                      AA 1+970529
                                          CA 1996-2237473 1996111E <--
     CA 1137473
                                      EP 1996-942767 19961116 <--
    EF 8640.1
                      Al 1+980916
        E: AT, BE, CH, DE, ES, FE, GB, IT, LI, NL
     JF 1000500584 T2 20000115 JP 19-7-519822 19901118 <--
                           19951110 <--
    _ ... | Swood | 1
| Web | 1996-0318471
| Web | 1 - 1 | 1
PRAI UN 1995-560601
                           19961118 <--
    Whole blood is mixed with a reticulocyte reagent system that has a
A.P.
    reticulcoyte staining reagent and a diluent reagent, used in combination.
     This mixt, is incubated at room temp, for between about 15 min to about 4
     h. The incubated mixt, is then analyzed and the light
     scattering properties of the cells are detected, collected,
     differentiated, and quantitated. Data gathering includes, at least,
     10.degree. and 90.degree. light-scatter detection.
ST
    blood reticulocyte differentiation identification stain reagent;
    light scattering reticulucyte differentiation identification
ΊΊ
    Algorithm
      Light scattering
     Fatiruloryte
     Staining, buclesidal
     Stains, biological
        (readent system and method for reticulocyte differentiation and
        adentification)
     591-65-9, Azure B 19:4-16-6, New Methylen: Blue 67566-77-3, Oxazine
ΙT
     750 86090-14-6, Brilliant Cresyl Blue
     FL: ARG (Analyt.cal reagent use); AMST (Analytical study); USES (Uses)
        (reagent system and method for reticulocyte differentiation and
        identification)
    62-76-0, Sodium oxalate 139-33-3, Disadium EDTA 1113-33-8, Ammonium
ΊT
    chalate 7447-40-7, Potussium chloride, analysis 7553-79-4, Dibasic stdrum phosphate 7647-14-5, Sodium chloride (NaCl), analysis
     7/7e-77-0, Monobasid botassium phosphate (00043-99-8, Potassium oxulato
     149 (3-0 -c, N-Tetraderyl-N,N-dimethyl-F-ammenic-l-propanesulfonaté
     55.365-84-4, Proulin 300 632.7-33-6, Dedecyl-.beta.-D-maltoside
     fo295-19-4, N-Dode:yl-N,N-dimethyl-9-ammonio-1-propanesulfonate
     RL: ARU (Analytical rate, unclassified); ANOT (Analytical study)
        (reagent system and method for retigulocyte differentiation and
        identification)
L79 ANSWER 20 OF 24 HCAPLUS COPYRIGHT 2002 ACC
    1995:95-693 HCAPLUS
2.1
EtM
     123:358026
ΤI
    Determination of particle concentration in suspension and
     apparatus thereof
     Yimanoe, Seijo
IN
     Cosmo Sogo Kenkyusho Ek, Japan; Cosmo Oil Co Ltd
EA
     Jon. Ebrai Tokkyo Koho, 7 pp.
\mathcal{E}\mathcal{O}
     CODEN: TEXXAF
DT
     Patent
     Japanese
LA
```

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IC
    FOM G01N021-49
    ICS G01N015-06
    70-6 (Inorganic Analytical Chemistry)
    Section cross-reference(s): 74
FAN.CHT 1
    PATENT NO.
                    KIND DATE
                                     APPLICATION NO. DATE
                                         -----
     ______
    TP 07198605 A3 19950801 TP 2820879 B1 19981105
                                         JP 1993-349459 19931228 <--
Ε·Ι
AB
    The title method, suited for use in colored suspension,
    comprises measuring scattered and transmitted light from the
    suspension.
ST
    suspension particle concn turbidimeter
    turbidimetry algorithm
T T
   Algorithm
    Juspensions
      Turbidimeters
      Turbidimetry
       (detn. of particle concn. in suspension and app. thereof)
L79 ANSWER 21 OF 24 HCAPLUS COPYRIGHT 2002 ACS
    1994:404517 HCAPLUD
DUI.
    121:4517
ΤI
    Method for determination of test readent concentration to avoid
    prozone phenomenon in turbidimetric immunoassay
    Makane, Hyekanu
III
    Chimadou Corp, Japan
\mathbb{P}A
    Jpn. Kekai Tokkyo Koho, 8 pp.
SO
    CODEN: JKKKAF
\mathbb{D}^{n}
    Patent
F_{i,\perp}
    Japanese
    ICH G01N035-00
7,~
     ICS G01N033-536; G01N033-543
CC
    9-10 (Biochemical Mothods)
FAN.CHT 1
    PATED, NO. KIND DATE APPLICATION NO. DATE
    FATENT NO. KIND DATE
    7F 0/109740 AZ 19946402
7F 3102100 BZ 20001023
                                         JP 1992-25:947 19920930 <-+
    The title method comprises (1) reacting test readent with stds., (2)
AB
    measuring the reactions at a 1st time point and a 2nd
    time point, (3) analyzing the data by linear segression, and (4)
    extrapelating and detq. the non-prozone phenomenon concn. region
    of the test reagent. The invention is a rapid method for detq. proper
     test reagent concn. to avoid prozone phenomenor and to out the
    post. The method is also appropriate for automatic
    turbidimetric immunoassay for antiger or antibody detn.
    turbidimetric immunoassay prozone phenomenon prevention; linear
    regression test reagent concn
ΙT
    Antibudies
     FL: ANT (Analyte); ANST (Analytical study)
        (detn. of, prevention of prozone phenomenon by two reaction
        time points measurement and linear regression anal. for detg.
        test reagent concn. in turbidimetric immunoassay
        for)
ΤТ
    Antigens
     kL: ANT 'Analyte); ANST (Analytical study)
        (deth. of, prevention of prozone phenomenon by two reaction
        time points measurement and linear regression for detq. test
        reagent concn. in turbidimetric immuniassay for)
ΙΤ
    Mathematics
        (equations, for linear regression anal., two reaction
        time points measurement and, for dety. test reagent
```

gitomer - 09 / 583891 concn. for turbidimetric immunoassay' ΙT Serelogical reaction (prodone, prevention of, two reaction time points measurement and linear regression for detg. test reagent concn. for, for turbidimetric inumunoassay) ΙΤ Statistics and Statistical analysis (regression, two reaction time points measurement and, for detr. of test reagent concn. for turbidimetric inmumoassay) ΙT Immurscassay (turbidimetric, prevention of prozone phenomenon in, two reaction time points measurement and linear regression for detq. test reagent concn. for) ΙT Immunicassay (turbidimetric, automated, prevention of prozone phenomenon in, two reaction time points measurement and linear regression for detg. test reagent concn. for) ANSWER 11 OF 24 HOAFLUS COPYRIGHT 1002 ACS 1994:184778 HCAPLUS MI 120:184778 Li: I TΙ Mathematical model of toxicity monitoring sensors incorporating microbial whole dells ΑIJ Hadgett, Barry G. D. Res. Cent., Univ. Luton, Luton/Beafordshire, LUI ELE, UK $\mathbb{C}\mathcal{S}$ Analyst (Cambridge, U. K.) (1994), 113(2), 197-201 CODEN: AMALAO; ISSN: 0003-2654 $\mathbb{D}^{r}\Gamma$ Journal LEEnalish CC4-1 (Toxi tology) A model is presented that describes aspects of the transient and A.H steady-state behavior of toxicity monitoring biosensors that incorporate living microbial cells immobilized in a thin layer between an amperometric electrode and a porous (nontertuous) membrane. In the example considered here, respiratory or photosynthetic electron-transport activity is menitored by using artificial redox negrators to divert electrons from the electron-transport systems to the working electrode goised at a suitable reducing potential. Such biosensors are being developed for a range of environmental monitoring applications. math. model is used to demonstrate how the response of practical devices can be manipulated and to indicate potential pitfalls in the interpretation of toxicity assessment data derived by such biosensors. math model toxicity analysis biosensor microbe STSimulation and Modeling, biological Ι'. (math., of toxisity manitoring minorbial biosensors) Ι". Texibity (meanitering of, by microbial biosensors, math. model of) ΙŢ Biosensors (microbial, amperometric, toxicity monitoring by, math. model ○f) L/9 AMEWER 13 OF 24 HOAPLUS CORVEIGHT 2002 ACS 1994:101289 HCAPLUS A11Dil I 120:101289 Process for the analytical determination of the concentration of Τi a component of a medical specimen Somaefer, Rainer; Berding, Christoph; Lang, Fridl; Eleider, Wilhelm; Wolf,

III Sanaefer, Rainer; Berding, Christoph; Peter

PA Boehringer Mannheim G.m.b.H., Germany SO Ear. Pat. Appl., 15 pp. CODEN: EPEMEDW

DT Patent

Là German

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ICM G01N033-53
IC
     ICS G01N021-47; G01N021-27
    9-10 (Biochemical Methods)
CC
FAN.CNT 1
                     KIND DATE
    PATENT NO.
                                          APPLICATION NO. DATE
    EP 576873 A. 19940105
                                          EP 1393-109189 13930608 -:--
PΤ
    EP 6/6879 A: 19940518
EP 6/6879 BI 19981011
    EP 176479
        R: AT, BE, CH, DE, DK, EC, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
    DE 4221807 AT 1 (94010)
                                          DE 1992-4221:07 19920703 <--
    I-E 4.1.1807
                      CL 10940714
    .. 172939
ES 21..5391
US 5420040
CP 67777
                     E 1.981115
T: 1.990116
                                          AT 1993-1091-9
                                                            19930608 -:--
                                          ES 1993-1091-9 19930608 <--
    T **990116 A 199505 kg CP (0167501 A 1997)
                                                            19930629 -:--
                                          US 1993-8400f
                     A. 1 (940614
1 (920703 K--
                                          JP 1993-164642
                                                           19930702 ::--
PRAI LE 1992-4001307

    Reaction of a medical sample with readents produces a time

    it)-geoendert change in a measurable parameter S, where the conci
     . Claf a component of interest in the sample is correlated to an input
    variable X derived from S(t) and the calibration curve X = f-1(C) is not
    monotomic, so that a value of X may correspond to .gtoreq.1 value of C. A
    discrimination algorithm is provided for correlating X with a
     unique value of C using multivariate statistics. The reaction
    may be a specific binding reaction, e.g. immunoppth., where S is
     turbidity. The method was applied to immunol, detr. of albumin in
     urine with a com. Rit by turbidimetry.
    turbidity immunoassay multivariate statistics; albumir.
    uring immunoassay turbidity
ΙT
    Urine analysis
        (a.bumin detn. in, by turbidimetric immunoassay,
        multivariate statistics in relation to)
ΙΤ
       (clin., specific binding assay using, multivariate statistics
        in)
    Ferritins
ΙΤ
     RL: ANT (Analyte); ANST (Analytical study)
        (detn. of, in blood serum by latex-enhanced turbidimetric
        immumoussay, multivariate statistics in relation to)
    Albumins, analysis
ΙT
     FL: ANT (Analyte); ANST Analytical study)
        (letn. of, in urine by turbidimetric immunbassay,
        multivariate statistics in relation to)
    Elook analysis
TT
        (ferritin detn. in, by latex-enhanced turbidimetric
        immuniassay, multivariate statistics in relation to)
TT
    Antibodies
     FL: ANST (Analytical study)
        (irmob: lized, for clim. immunopptm. assay, multivariate
        statistics in relation to)
     Receptors
     FL: ECT (Emastant)
        (reaction of, with ligands in turbidimetric clin. anal.,
        multivariate statistics in relation to)
TΤ
     Ligands
     EL: ROT (Reactant)
        treaction of, with receptors in turbidimetric clin. anal.,
        multivariate statistics in relation to)
     Statistics and Statistical analysis
ΤT
        .discriminant, in turbidimetric specific binding
        ascay, in clin. anal.)
ΙT
     Immunoassay
        jimmunopptn., immobilized antibody for, in clin. anal.,
```

multivariate statistics in relation to)

```
ΙΤ
     Statistics and Statistical analysis
        (multivariate, in turbidimetric specific binding
        assay, in clim. Anal.)
L79 ANSWER .4 OF 24 HCAPLUS COPYRIGHT 2002 ACS
   1983:61103: HCAPLES
A:1
ECL
    99:31:0:1
TI
    Analysis of Iq
    Shimadzu Corp., Japan
F A
    Jpn. Kakai Takkya Kaha, 4 pp.
\leq \omega
    CODEN: JEKKAF
\Gamma \cdot \Gamma
   Patent
LJ_{i}
   Japanese
    G01N033-54
I \cap
("t"
    15-1 (Immunochemistry)
FAN.CNT 1
                                    APPLICATION NO. DATE
    FATENT NO.
                    HIND DATE
    JP 58113758 AJ 19830706 JP 1981-211308 19811226 <--
E- :
    The detr. of Iqs in blood serum using immunoturbidimetry is
     improved by using sample blank channels in addn. to conventionally used
     mal. charmels for the measurement of the turbidity absorption
     ut 340 mm. The parameters for the measurement are defined, and
     math. formula for the calcn. of Ig concns. are
     presented. Accurate Ig concns. were detd. even in the presence
     of 62.5-150 mg Hb/dL or 2.5{	ext{-}}10.0 mg bilirubin/dL is blood serum.
ST
     Iq detr immunoturbidimetry
IT.
    Blood analysis
        (I) detn. in human, by immunoturbidimetry)
Ι".
     Immunoglabulins
     FL: ANT (Analyte); ANST (Analytical study)
        (detr. of, of blood of human, by immunoturbidimetry)
ΙT
     Immunectemical analysis
        (immunoturbidimetry, of Ig of human)
= - d 185 all tot
185 ANSWER 1 OF 10 HOAFLUS COEYRIGHT 2002 ACS
A:::
    -2002:125572 HCAFLUS
    136:139897
    Device for monitoring liquid biological medium
    Vašilevskil, A. M.; Kornilov, N. V.
Ι::
PΣ
    Lugara
30
    Fuss., No pp. given
    CODEN: RUKEE7
\mathbb{D}^{n}:
    Patent.
L.S.
    Folsenar.
Ir.
    HUM G01N021-31
     + ?= 7 (Pharmaceuticals)
Cr.
     Section cross-reference(s): 9
FAN.CHT 1
    PATENT NO. KIND DATE APPLICATION NO. DATE
    EM 2161791 CL 20010110 RU 1993-123692 19981230 <--
PΙ
    Monitoring of liq. Hool. medium, e.g., components of dialysis liqs. in
An
     hemodialysis is based on the formation of light beam of a
     montinuous spectrum source in a controlled zone of lig. biol. medium.
     Change of characteristics in this section is detd. and spectral coeffs. of
     correlation of absorption dynamics per each analyzed domponent are
     computed. Later, luminous flux is transmitted through a dish with
     liq. bil. medium, radiation passed in spectrum is decompd. and
```

```
transmission coeff. of liq. medium is measured. Finally concn.
     of analyzed components is computed by spectral coeffs. of
      tirrelation if absorption dynamics. The proposed device has a
     light source, optical system forming light beam, a dush
     with bicl. liq. medium flowing through it, a spectrometer and controller
     installed in series, controlling computer, unit controlling
     parameters of repording and tuning of spectrometer, a processor of the
     spectra, unit controlling monitoring parameters, timer, input data,
     processing, output data and display units, and unit of algorithms
     . Controlling computer is connected to controller on one side
     and to unit matrolling parameters of recording and funing of spectrometer
     and import data unit on the other side. The latter is connected in its
     turn to unit controlling monitoring parameters. First output of
      controlling computer is demonstred to the input of processor if
     present spectrum whose output is linked to display. Second output of
     rentrelling computer is dennected to timer, processing unit and
     unit of algorithms connected in series. The unit of
     algorithms is connected to display and cutput data unit. The
     catput of processor of present spectrum is connected to the input of
     rimer.
     liq biol medium device
ST
ΙΤ
     Algorithm
       Computer application
       Light
         (device for monitoring liq. biod. medium)
ΙT
     Dialysis
         (hemodialysis; device for monitoring liq. biol. medium)
     ANSWER 2 OF 10 HUAPLUS COPYRIGHT 2:02 ACS
L25
     L001:439732 HCAFINS
A:1
DN
     198:78958
     Method and apparatus for controlling the manufacturing quality of a moving
T 1
III
     Workman, Jerome J., Jr.
     Fimberly-Clark Worldwide, Inc., USA
PA.
     FCT Int. Appl., 27 pp.
SO
     CODEN: PIXXD2
D"
     Hatent
L÷.
     Englash
I \in
     ICM G01N021-86
     ICS G01N021-31; E21H023-73
     48-10 (Thit Operations and Processes)
     Gestion cross-reference(s): 74
FAN.CNT 1
     भवासाम अत
                        MIND DAME
                                              ASSLICATION NO. DATE
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                                                ______
                                         WO 2000-U334630 20001220 ≪--
                       Al 20010705
PΙ
     W0 2 01045462
         W: AE, AG, AL, AM, AT, AU, AL, BA, BE, BG, BE, BY, BZ, CA, CH, CN, CE, CU, CZ, DE, EK, EM, DG, EE, ES, FI, GR, GD, GE, GH, GM, HR, HU, 1D, IL, IN, IC, JE, FE, KG, FE, FE, KJ, LC, LE, LE, LS, LT, LG, LV, MA, MD, MG, ME, MI, MW, MK, MZ, NO, NZ, PL, ET, RO, RU,
              SD, SE, SG, SI, SK, SL, TJ, SM, TF, TT, TC, MA, UG, UZ, VN, YU,
          ZA, CW, AM, AZ, BY, FG, ED, MD, FY, TT, TH

EW: GH, GM, FE, LS, MW, MZ, CD, JL, EZ, TZ, CG, CW, AT, BE, CH, CY,

DE, DE, ES, FI, FE, GB, GE, IE, IT, LU, MC, NJ, PT, SE, TR, BF,
              BJ, CE, CH, CH, CM, CA, CH, CW, ML, ME, ME, JN, TD, TG
474713 A 199911.9 ---
PRAI US 1999-4747_9 A 199910.9 --- AB A method and app. are disclosed for detecting the compn. of a moving web
     product on a real-time madis during the ranufg, process.
     "pestrometric monitoring equipment operates to derive information
     regarding phys. and/or chem. properties if the web at multiple
      locations in the web's cross direction. Data from a plurality
     of spectral regions can be dombined to produce a vector contg. accurate
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IE spectroscopy

information regarding the web's compn. This information is derived using multivariate math. techniques to yield a spatial data matrix for each component of interest. Compn. information contained in the spatial data matrix can be reprojected as a "virtual componinap," or compared against ideal profiles stored in a computer memory. controlling manufg quality moving web; app controlling manufg quality moving web Apparatus ΙΤ Optical detectors Quality control (method and app. for controlling manufg. quality of moving web) RE.CNT THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD F.E (1) Abk Ind Systems Inc; EP 0081188 A 1995 (2) Qualico Gmbh; DE 19709963 A 1998 (3) Siemens Aq: DE 19683477 A 1998 HCAPLUS (4) Siemens Aq; DE 19830328 A 1999 LH5 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2002 AGS 2001:168235 HCAPLUS AM134:204739 DIL Method for determination of tissue analytes using MIR, adjacent visible srectrum and discrete NIE wavelengths 111Scedina, Thomas G.; Fawluczyk, Romuald; Cadell, Theodore E. CME Telemetrix Inc., Can. ĒΆ SO FCT Int. Appl., 33 pp. CODEN: PIKKD2 $\mathbb{D}\mathbb{T}$ Eatent Er.qlish HTM G01N021-35 ICS G01N021-31; A61B005-00 CC 9-9 Brochemical Methods: FAN.CET 1 FATENT NO. APPLICATION NO. DATE HIND DATE ____ _____ -----WO 2001016577 Al 20010708 - WO 2006-CA1000 20000931 ∹--W: CA, CE, TE FW: AT, BE, CH, CY, DE, DK, EC, FI, FR, GB, GR, IE, IT, LC, MC, NL, PT, JE E14577 A1 16626619 EP 2001-985344 21000831 H--R: AT, BE, CH, DE, DK, ES, FP, GP, GR, IT, LI, 1U, NL, SE, MC, PT, EF 1214577 IE, SI, LT, LV, FI, FO, ME, CY, AL PRAI US 1999-1515:7P E 19990831 We 2000-CA1000 W 20000831 Described is a method for measuring the concn. of a blood AB constituent within a body part (80) of a living subject which comprises irradiating a body part of the subject with a continuum of a broad spectrum of radiation in adjacent and near IR range of the electromagnetic spectrum; collecting the band of radiation after the radiation has been directed onto the part; dispersing the continuum of collected radiation into a dispersed spectrum of component wavelengths onto a detector (120), the detector taking measurements of at least one of transmitted or reflected radiation from the collected radiation; and transferring the measurements to a processor (300), and then measuring the same kind of absorbance or reflectance with respect to one or more discrete wavelengths of radiation from the longer near IR range and using the measurements to calc. the concn. of the constituent. tissue analyte NIE adjament visible spectrum spectroscopy ST (finger; tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIE wavelengths)

Optical detectors

```
(near-IE; tissue analytes detn. using NIR and adjacent visible spectrum
        and discrete NIE wavelengths)
TΤ
    Blood analysis
        (noninvasive; tissue analytes detn. using NIE and adjacent visible
        spectrum, and discrete NIE wavelengths)
ŦΤ
    Algorithm
    Animal tissue
     Body, anatomical
      Computers
      Optical detectors
      Spectrometers
     Spectroscopy
        (tissue analytes detr. using NIR and adjacent visible spectrum and
        discrete NIR wavelengths)
ΙT
     50-99-7, D-Glucosé, analysis
     EL: ANT (Analyte); ANST (Analytical study)
        (tissue analytes detr. using NIE and adjacent visible spectrum and
        discrete NIR wavelengths)
P.E. CNT 4
            THERE ARE 4 CITEL PEPERENCES AVAILABLE FOR THIS RECORD
(1) Domian, G; WO 9637259 A 1996
(2) Guthermann, H; US 5818048 A 1998 HCAPLUS
(3) Khalil, G: US 5747806 A 1998 HCAPLUS
(4) Lepper, J; US 5743262 A 1998
LES ANSWER 4 OF 16 HOAPLIS COFFEIGHT 2002 ACS
A11
   1999:811438 HCAPLUS
11:1
   192:47332
ΤI
   A method in quality control of a spectrophotometer
I!I
   Hansen, Heine
F.A.
   - Radiometer Medical A'S, Den.
SO
   FOT Int. Appl., 61 pp.
    CODEN: PIXXDO
ľΤ
    Patent
    English
LA
    ICM G01N021-31
11
    IGS G01N033-49; G01J003-42
CiG
    9-5 (Baschemical Methods)
    Section cross-reference(s): 33
FAN.CNT 1
                                         APPLICATION NO. DATE
     PATENT NO.
                 HIND DATE
                     Al 19991223
                                                           19990610 <--
    WO 3466310
                                          WO 1999-DB313
        W: JE, US
        FW: AT, BE, CH, CY, DE, DE, EJ, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, JE
                      A1 20010318
                                          EP 1999-924801 19990610 4--
     EP 1086166
        JP 2002518670
                                          JP 2000-555079
                                                          19990610 <--
PHAI DH 1998-783
                           19950612
                      F_{\lambda}
                     W
                          19990610 ----
     WO 1939-DKF13
    Methods for calibration of spectrophotometers, esp. oximeters for blood
     anal., are described which entail using the spectrophotometer to det. a
    spectrum Am(.lambda.) of a fluid ref. sample contg. a dye, and detg. a wavelength shift .DELTA..lambda. :rom
     C.DELTA..lambda.(.lambda.)Am(.lambda.), where C.DELTA..lambda.(.lambda.)
     is a predetd, coeff, vector previously stored in a memory of the
     spectrophotometer. Vectors for interferences (e.g., fetal Hb) may also be
     stored and used to produce calcd. spectra for which the effects
     of the interference are minimized. Spectrophotometers provided with
     memory with a math. parameter for the detn. of a
     wavelength shift of the spectrophotometer, and a processor that is
```

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connected to the memory and that is adapted to calc. the
    wavelength shift AX from the math. parameter and from a
    spectrum detd, with the spectrophotometer on a dye-contg, ref. sample are
    also esscribed. Alternately, a lamp with a known emission spectrum may be
    assection place of a ref. sample. Spectrophotometers may be prepd. for
    mallbration by detg. a first ref. spectrum of a ref. sample contg. a dye
    in a first concn. with a ref. spectrophotometer, detq. a first
    derive of the first ref. spectrum, and detg. from at least the first ref.
    spectrum and the first deriv. a math. parameter from which a
    wavelength shift of the spectrophotometer can be detd., and
    storing the math. parameter in a memory of the
    spectrophotometer.
    spectrophotometer calibration; oximeter calibration
    Calibration
      Spectrometers
        (malibration of spectrophotometers and spectrophotometers equiped for
        the calibration)
    Analytical apparatus
      Analytical apparatus
    Medical equipment
    Medical equipment
        (ownmeters; calibration of spectrophotometers and spectrophotometers
        equiped for the calibration
    3520-41-1, Solforhodamine B
    FL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
        (valibration of spectrophotometers and spectrophotometers equiped for
        the calibration)
    #034-03-3, Fetal hemoglokin
    EL: ARY (Analytical role, unclassified); OCU (Occurrence, unclassified);
    ANST (Analytical study); OCCU (Coburrence)
        1941 ibration of spectrophotometers and spectrophotometers equiped for
        the calibration:
             THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
PE.CNT 4
(1) Abbott Laboratories; EP 0167816 A2 1986
(D) Asland Oil, Ind; WO 0408225 Al 1994 HCAPLUS
(a) Atsuhire, I; 00 5592291 A 1997 HCAPLUS
(4) Ciba Corning Diagnostics Corp; WO 3630742 Al 1996 HCAPLUS
185 ANSWER 5 OF 10 HCAPLUS CONTRIGHT 2002 ACS
    1994:307860 HCAPLUS
     130:303767
     Three wavelength in-vivo analyte detector
    Dopson, Peter J.; Turner, Socit J.
Abbott Jahorstonies, USA
    Frit. TK Pat. Appl., 18 pp.
    PRODEN: BAKKDU
    Patent
   English
    ICM G01N021-31
ICA A018005-00; G01N001-30
    e-l (Biochemical Methods)
FAN.CHT 1
    PATENT NO. KIND DATE
                                         APPLICATION NO. DATE
     #B UNU8279
                    A1 14990217
                                          GB 1997-17134
                                                           13970812 <--
                                           JP 2000-506877
     TP 2001513351 T2 20010904
                                                           13980805 <--
PEAT 3B 1997-17134
                           19970612
                     А
    W() = 1 \cong (m + (GB, (G, S))) W
                           19930605
    A device for measuring the concn. of an analyte in blood in vivo
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... Hisphosed. The device comprises (1) a transmitter for illuminating a body part with light at a plurality of predetd. wavelengths; (2) a detector for receiving light from the

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body part and generating input signals representative of the intensity of
    received light at each of the wavelengths; and (?) a
    computer coupled to the detector for generating an output signal
    representative of the conc. of analyte in the blood in the body
    part by anal. of input signals received from the detector. The
    transmitted or reflected intensity of light at three discrete
    wavelengths is analyzed by computer. The analyte is
    esp. glucose and the body part is a finger. A formula is given for
    calcg. the cutput signal from light received at three
    discrete wavelengths.
     in vivo analyte detector; blood glucose in vivo light detector
    Hand
        finger, blood glucose detn. in; three wavelength in-vivo
       analyte detector)
    Mathematical methods
        efer calcq. output signal from light received at
        three discrete wavelengths; three wavelength
        in-vive analyte detectio)
    Analytical apparatus
    Pleed analysis
    Fody, anatomical
      Computers
      Optical fibers
     Ehstadindes
      Sensors
        three wavelength in-vivo analyte detector
     50-99-7, Gludose, analysis
    EL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL
     (Biological study); USES (Uses)
        (as analyte; three wavelength in-vivo analyte detector)
    50-94-7, D-Glucose, biological studies
    FL: Red (Biological occurrence); BSU (Biological study, unclassified);
    FIGL (Biological study); OCCU (Occurrence)
        (blood; detn. of; three wavelength in-vivo analyte detector)
L85 ANSWER 6 OF 10 HCAPLUS COFFRIGHT 2002 ACS
    1999:267960 HCAPLUS
    130:264412
    Infrared multi-wavelength non-invasive measurement of
    kload companent concentrations
    Amerov, Airat K.; Jeon, Kye-Jin; Kim, Yeon-Joo; Yoon, Gil-Won
   Samsung Electronics Co. Limited, C. Korea
    Brit. UK Fat. Appl., 27 pp.
    CODEN: BAKKDU
    faters
    Englian
    ICM A618005-00
     ICS G01N021-31; G01N021-35
    9-1 (Biochemical Methods)
     Section subserveference(s): 1
FAN.CNT 1
                 KIND DATE
                                          APPLICATION NO. DATE
     PATENT NO.
                           _____
                                          ______
                     ____
    GB 15.73015
                      Al 14990:10
                                          GB 1998-18315
                                                          19980821 <--
                     B2
    GB 1333015
                           20020.13
                     Al 1.090312
     FR 1.769043
                                          FF. 1998-11103
                                                         19930904 <--
    FF. .17+6043
                     31 200000F11
                     А
PRAI EE 1997-45970
                          19470909
                     А
     EE. 1398-21963
                          1 +930+012
    A method and device for moninvasive measurement of plood component
    concns. utilizes pulsed polychromatic light source
     emitting in particular light in the near IR range 810-1850 nm.
```

The light is back scattered from or transmitted through a part

```
of a patient's body. Back scattered light from blood-contg.
     tissues and blood vessels has information on blood component
    concns. That light is properly collected to avoid the
    surface reflection from the skin surface and provide minimization of the
    effects of changes in the scattering background. The concns. of
    blood components is calcd. from the spectral anal. based on
    selected wavelengths by a proposed algorithm. A
    microprocessor calcs. a ratio and dets. the blood component
    concis. by comparing the ratio with a calibration curve stored in
    a memory of the microprocessor. The calcd. concn.
    values can be used for clin. use or for a home test.
ST
    IF multi wavelength noninvasive blood component
ΙT
    Algorithm
     Blood analysis
    Blood vassel
       IR spectrometers
       Light sources
       Optical detectors
     Pharmaceutical analysis
     Skin
        (IF multi-wavelength non-invasive measurement of
        blood component concns.)
    Albumins, analysis
TΤ
    Hemoglobins
    FL: ANT (Analyte); ANST (Analytical study)
        (IR multi-wavelength non-invasive measurement of
        blood component concns.)
ΙΤ
     Electric circuits
        (analog-digital converters; IR multi-wavelength
        non-invasive measurement of blood component concns.)
ΙT
     Computers
       (microprocessors; IR multi-wavelength non-invasive
       measurement of blood component concns.)
     IF. speatrescopy
TΤ
        (near-IR; IR multi-wavelength nin-invasive
        measurement of blood component concns.)
     50-99-7, Glucose, analysis 57-98-5, Cholesterol, analysis 64-17-5,
ΙT
     Ethanol, analysis
     EL: ANT (Analyte); ANST (Analytical study)
        (IE multi-wavelength non-invasive measurement of
        klased dempenent concns.)
L35 ANSWER / OF 10 HCAPLUS COPYRIGHT 2002 ACS
    1998:696989 HCAPLUS
AH
DH
    119-187556
T.
    Spectrophotometric analysis for nemoglobin analysis in blood
    Jarman, Hoyer Kristin; Pologe, Jonas A.
I:1
    Ohmeda Inc., USA
FΑ
    Eur. Bat. Appl., 13 pp.
SO
    CODEN: EPKKDW
DΤ
    Patent
    English
LA
     ICM G01N021-31
TO
     103 A61B005-026; G01N033-487
CC
     9-5 (Biochemical Methods)
     Section cross-reference(s): 0, 13
FAN.CNT 1
                    KIND DATE
                                          APPLICATION NO. DATE
     PATENT NO.
                           ______
                     ____
                                           EP 1998-302120 19980320 <--
                      A2 19981014
    EP 8710.16
                      A3 19990113
     EP 871026
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
```

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JP 10318915
                      ΑŌ
                         13331.04
                                          JP 1998-93431
                                                          19980406 <--
PRAI US 1997--35289
                          19970409 <--
   A two-stage statistical calibration and measurement method and system is
    disclosed for performing photoplethysmig, measurement of blood analyte
    concns. Concns. in a tissue sample of MetHb, 02Hb, Hb
    and CoHb are estd, by first estq. a concn. of MetHb (in a first
    stage) and subsequently, if the concn. of MetHb is within a
    predetd, range, then the estd. concn. of MetHb is assumed to be
    about the and this estd. concn. of MetHk is utilized as a "known
    value" in detg. the concns. If the remaining analytes O2Hb, Hb
    and COHb (in a second stage). By eliminating one "unknown" from the
    system of equations, these remaining analytes can be
    calcd. with increased accuracy. Each stage is performed using
    data obtained by transmitting light through the tissue sample
     (type fally a finger or earlebe). The transmitted light is
    demerated by four discrete light emitters, each emitter having a
    distinct spectral content.
    blood Hb analysis photoplethysmog spectrophotométry
ST
    Hemoglobins
ΙΤ
    FL: ANT (Analyte); BSU (Biol.gical study, unclassified); ANST (Analytical
    study); BIOL (Biological study)
        (markexyhemoglebins; spectrophotometric anal. for Hb anal. in blood)
I"
       (priotoplethysmag, analyzer; spectrophotometric anal, for Hb anal, in
       blood)
    Algorithm
ΙT
    Animal tissue
    Billiocea
    Plood analysis
      Mathematical methods
      Spectrometers
      Spectrophotometry
        (spectrophotometric anal. for Hb anal. in blood)
    Hemoglickins
ΙT
    Hemoglobins, methemoglobins
    Homoglobins, oxyhemoglobins
    FL: ANT (Analyte); BSU (Bitligical study, unclassified); ANST (Analytical
    study; ; bICL (Biological study:
        (opectrophotometric anal. for Hb anal. in blood)
Las Answer a of 10 HCAPLUS COPYRIGHT 2002 ACS
    1997:L51014 HCAPLUS
AE
    126:2:5573
DI:
    Method of quantitatively determining one or more characteristics of a
т:
    substance
III
    Evans, Feter Dilwyn; Barnett, Nicholas
    Johnson & Johnson Medical, Inc., USA
PA
    Eur. Fat. Appl., 13 pp.
EO
    GODEN: EEKKIW
DΤ
    Patent
L\Delta
    English
    ICM G01N021-31
I+1
    ICS G01N021-47; A61B005-30
    9-1 (Riochemical Methods)
    Section pross-reference(s): 13, 73
FAN.CNT 1
                 KIND DATE
                                          APPLICATION NO. DATE
    PATENT NO.
                                          -----
                           19970-05
    EP 760476
                                          EP 1996-306179 19965823 <--
                 I_{\lambda}:
                           19980004
     EP 760476
        A: DE, FE, GB, SE
PRAI GB 1995-17366 19950-24
                                     < --
   A method is described for quant, detg. .gtoreq.1 characteristic of a
```

substance by near-IR spectroscopy, wherein the characteristic is, e.g., the HE or cytochrome concn. of a tissue of the numan or animal The method involves: irradiating a point of the substance with あるます. radiation at .gtoreq.2 distinct wavelengths, measuring the intensity of the radiation detected at 2 locations, detg. the optical path Tengths of the radiation between the irradn. point and the 2 detecting locations, and detg. the effect of the divergence of the radiation reaching the 2 locations. The relative coupling efficiencies of the 2 detectors are detd. by the use of a 2nd emission point equidistant from the 2 detectors. The characteristic being measured is then detd, by the intensity of the radiation detected at the detecting locations with the result modified by accounting for the optical path lengths to the detecting locations, the detector coupling efficiencies, and the effect of divergence of the radiation before reaching the detecting locations. An example shows a qual-channel sensor placed on the surface of the human nead for use in analyzing the perebral portex, but the invention also can re used to monitor noninvasively tissue Hp concn. in other parts of the body and may be useful in fields outer, as plastic surgery and vascular surgery. pady tissue analysis near IR spectroscopy; Hb deth derebral cortox IR detector Brain (perebral cortex; quant. anal. of body tissues by near-IE spectroscopy) IR spectroscopy Optical detectors (mear-IR; quant. anal. of bedy tissues by near-IR spectroscopy) Surgery (plastic; quant. anal. of body tissues by near-IE spectroscopy) Amimal tissue Body, anatomical Electroluminescent devices Mathematical methods -guart. anal. of body tissues by near-IF spectroscopy) Cyttechalomes Hemoglobins FL: ANT (Analyte); ANST (Analytical study) ,quant. anal. of body tissues by near-IE spectroscopy) Flood vessel Eldod vessel (surgery; quant. anal. of body tissues by near-IR spectroscopy) Curmery Surgery (vascular; quant. anal. of body tissues by near-IR spectroscopy) 77%1-18-8, Water, processes FL: FEF (Physical, engineering or chemical process): PROC (Procous) equant. anal. of body tissues by near-IR spectroscopy) LB5 ANGWER 9 OF 10 HCAPLUS COPYRIGHT 2002 ACS 1990:422863 HCAPLUS 117:21863 Thetemeter-based apparatus for noninvasive determination of total hemoglabins in blood Hamaguri, Kenji Minolta Camera E. K., Japan Jpn. Kosai Tokkyo Kohe, 6 pp. CODEN: TEXXAF Patent Japanese LCM AG1B005-14 ICS G01N021-31

ST

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PA

SI)

DT.

LA

IC

CC

FAN.CHT 1

PATENT NO.

0-1 (Biochemical Methods)

KIND DATE

APPLICATION NO. DATE

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_______
    JP 04040940 A2 19920212 JP 1990-149527 19900607 <--
PΙ
    The title app. consists of a device to irradiate a test subject with
AΒ
    light having .gtoreq.2 different wavelengths that show
    different apsorbance seeffs, for Hbs and water, a light receiver
     for the permeated or reflected lights with different
    wavelengths, and a device for calcg. total Hos. in a
    blood sample based on the ratio of the pulsating components of the outputs
     corresponding to the various wavelengths. The total Hb contents
     can be calcd. using the equation: total Hb
    concn. = a E2 + b [a, b = integral no; E = the ratio of the
    pulsating components]. The method is moninvasive. Diagrammatic views of
     the app. are presented.
ST
    app photometer noninvasive Hb detn; math equation
    total Hb detn
ΙT
    Hemoglobins
    EL: ANST (Analytical study)
       (detr. of fotal, nominvasive, photometer-based app. for)
IΤ
    Photometers
       (in app. fer noninvasive total Hb detn. in blood)
ΙT
    Blood analysis
       (total Hb detn. in, nominvasive, photometer-based app. for)
ΙT
    Mathematics
       (equations, for total Hb detn. with photometer-based app.)
L85 ANSWER 10 OF 10 HCAPLUS COPYRIGHT 2002 ACS
    1992:169643 HCAPINUS
AN
[11]
    116:169643
TT
   Fetermination of hemoglobin oxygen saturation in erythrocytes for
    circulation dynamic menitoring
III
   Ishikawa, Muneharu; Yamamoto, Tetsuya; Yanebako, Makoto
FA
   - Mowa K. K., Japan
SO
    John. Rohai Torkyo Koho, 4 pp.
    CODEN: JEKKAF
Γ Τ
   Fatent
LA
   Japanese
    ICM A61B005-14
IC
    ICS G01N021-31
CC
    (HS (Picchemical Methods)
FAN.CMT 1
    PATENT NO. KIND DATE
                                       APPLICATION NO. DATE
    ΕI
    The title method involves: irradn. of test subjects with multiple
A.B
    interfering light having different wavelengths,
    eliminating the scattered light having frequency components
    equiv. to the blood flow, and detg. and comparing other frequency
    components to det. the Hb sath. in erythrocytes for monitoring of the
    blood circulation dynamics. The method is accurate. Equations
    for the calcn. are presented.
    Hb oxygen sath erythrocyte directlation dynamic; spectrometry Hb oxygen
ST
    blood circulation
ΙΤ
    Circulation
       (dynamics, spectrometric detr. of oxygen sath. in erythrocyte for)
    Hemoglobuns
    EL: ANST (Analytical study)
       (sath. of, in erythropyte, spectrometric detr. of, for monitoring
       circulation dynamics)
ΙT
    Mathematics
       (equations, for erythrocyte Hb satn. detn. for monitoring
       circulation dynamics)
    0782-44-7, Oxygen, analysis
ΙT
    FL: ANST (Analytical study)
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(erythrocyte Hb sath with, detn. of, for moditoring circulation dynamics)

=> fil wpix FILE 'WPIM' ENTERED AT 08:01:46 ON 08 JUL 2002 COPYRIGHT (C) 2001 THOMSON DEFWELT <21020 C4 UP> FILE LAST UPDATED: 04 JUL 1902 MOST RECENT DEPWENT UPDATE .:CO.4. <20+142/EW> DEPWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE (b) Update 1001-42 does not contain any new polymer indexing <<</p> Not The BATCE option for structure searches has been enabled in WEINDEN/WEIDS and WEIK Door 👉 : PATENT IMAGES AVAILABLE FOR PRINT AND LISPLAY 🐵 🖘 Not FOR DETAILS OF THE PATENTS CONERED IN CURRENT STRATES, SEE littp://www.derwort.com/dwpi/updates/dwpicov.index.html <<< · OF FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE, PLEAME VISIT: http://www.stn-international.de/training_center/pytents/stn_guide.pdf <<< 1989 FOR INFORMATION ON ALL DEFWENT WORLD PATENTS INDEX USER GUIDEC, PLEASE VISIT: http://www.derwent.com/userquides/dwpi guide.html <<< so id all abed tech tot 1119 L110 ANSWER 1 OF LO WEIK (C) \$802 THOMSON DERMENT 7002-077015 [11] WPIK - bnc c.0002-0131 1 DMM N2002-056943 Analysis of samples, especially for minimum inhibitory concentration determination, comprises mathematically combining spectral data to provide at least two drowth indicator values. -11-BU4 **SO3** CLYDE, M; OCONNELL, M A; PARMIGIANI, G; 111 TURNER, D J; WILES, T M; O'CONNELL, M A ĪΑ BECT) BECTON DICKINSON & CO CTC 2: 160564 A2 2001120% (200211)* EN 32p G01N021-51 K--E: AL AT BE CH OY DE DW ES EL FR OB DE LE LT LL LT LU LV MC MK NL PT MF 1160564 RO SE SI TR 2:010526; JP 2002125637 A JP 2001-160535 20010529 PRAI US 3000-583891 20000531 ICM C12Q001-02; C12Q001-18; G01N021-31 C12M001-34; G01N021-25; G01N021-64; G01N021-78; G01N033-15; G01N033-48; G01N033-483; G06F019-00 EP 1160564 A UPAB: 20020215 ĂВ NOVELTY - Method (A) for analyzing a sample comprises: (a) directing different analyzing Light wavelengths onto the sample in a sample well; (b) detecting a resultant light wavelength emanating from the sample for each analyzing light wavelength; (c) generating a result value representative of each resultant light wavelength; and

(d) mathematically combining the result values to provide growth indicator values.

DETAILED DESCRIPTION - Method (A) for analyzing a sample comprises:

- (a) directing different analyzing light wavelengths onto the sample in a sample well;
- (b) detecting a resultant light wavelength emanating from the sample for each analyzing light wavelength;
- (c) generating a result value representative of each resultant light wavelength; and
- (d) mathematically communing the result values to provide at least two growth indicator values, each representing a growth characteristic of the sample.

INDEPENDENT CLAIMS are also included for the following:

- (1) a method (B) for determining a minimum inhibitory concentration (MIC) value for a sample in a container having several wells, each containing a portion of the sample and a growth-affecting material, which comprises taking a set of readings for each well at a series of time intervals to provide a set of values for each well at each time interval, mathematically combining the sets of values for each well to provide a characteristic value for each well, drouping the characteristic values into droups representative of groups of wells and comparing the characteristic values with each other in each group to determine a MIC value for each group of wells;
- (L) a computer-regulable meaium with instructions for performing the operations of method (A), and
- (3) a computer-readable medium with instructions for performing the operations of method (B).

USE - The method is useful for analyzing the antibiotic suspentibility of biological samples and for determining the minimum inhibitory concentration (MIC) values of antimiorobial materials.

ADVANTAGE - The method uses multiple drowth indicators to provide increased accuracy and integrity of results.

Dwg. 0712 OPÎ EFI

FS

FA

cg: B11-c0/EC; B11-c08B; B11-c08C; B11-c09; B17-K04A4; B14-A01 MC: EPI: SUB-E04A5

UPTK: 23300218 TECH

TECHNOLOGY FOCUS - BIOLOGY - Breferred Method: The sample is contained in a series of sample wells and steps (a)-(d) are performed on each well at a series of time intervals so that step (d) provides a set of growth indicator values for each well at each time interval.

The method also comprises mathematically combuning certain values in the sets of growth indicator values for each well to provide a characteristic value itr each well, optionally grouping the undractedistic values into group's representative of groups of wells and comparing the characteristic values with each other in each group to determine in which wells within each group sample growth is inhibited.

```
L110 ANSWER 2 OF 10 WEIK (C) 2002 THOMSON DERWENT
```

2002-050099 [97] WEIK ΑN

CF: 2001-627611 [69]

INN N2002-036924

ΊI Method and device for analysis of substance mixtures using spectral analysis employs binary filters.

Įα S03

IMEPPICH, B; MUELLEE, G

*BECT) BECTON DICKINSON & CO; (LASE-11) LASER & MEDIZIN ΞÆ TECHNOLOGIE GMBH

CYC

A1 2001101a (200207)* DE 10018940 13p G01NUR1-25 FΙ AU 2001(65860 A 20011030 (200219) G01N021-31

ADT DE 10018940 A1 DE 2000-10018940 20000417; AU 2001065863 A AU 2001-65863

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20010406
    AT .001065863 A Based on W0 100179815
PPAI 05 L000-10016040 20000417; DE L000-10018941 20000417
    ICM G01H0.1-.5; G01N021-31
     183 4013003-83
ΑF
    DE 1001-40 A UPAB: 1002(10)
    MOVELTY - The aspectral power P(l) of the substance mixture (1) is split
     into M part beams by a beam applitter (?). After baseing through the
     spectral filters (3) the remaining placer is determined by broad band
    detectors (4). They are expertially bonary filters with a 3 or 1 output
     depending on the wavelength. The determination of the spectral part beams
     transmitted is by using the algorithm described in the patent.
          UNE - To detect changes in substance mixtures.
          ALMANTAGE - With the use of binary filters changes in the substances
     produce maximum changes in the signal vectors formed from the signals from
     the individual detectors.
          DESCRIPTION OF DRAWING(S) - The figure phoos a block diagram of a
     metroid to the present invention.
          Substance mixture 1
     Beam splitter 3
         Apectral filters :
     Fetedtors 4
         Spectral power P(1)
     Fwa.176
\mathbb{F}\mathbb{C}
    EFI
ΞA
     AP; GI
Mr.
     EFI: SOB-EC4A
L110 ADSWER E OF 2: WEIN (1) . 002 THOMSON DERWENT
    2001-027011 [75]
                       WPIH
11A
\mathbb{C}F
     2002-050099 [70]
DEN NL001-467412
                        DMC C1001-187018
    Multiple filter photometer used for determining small denoentration
Τi
     changes in a multiple component mixture comprises wide hand detectors for
     receiving the transmitted or remitted radiation with filters.
    √04 s03
Dit.
I.:
    ELPICH, B; MUELLER, G
     (BECT) BECTON DICKINSON & CO; (LASE-U. LASER & MEDICINI
PΑ
     TECHNOLOGIE GMEH
CIC
    .15
     DE 10018941 Al 2001101: (20017:)*
                                                      GUINDH1-25
PI
                                               12p
     Wo 1001070815 A1 20011025 (20017:) EN
                                                      G01N021-31
        RW: AT BE CHICY DE DK EA ES FI FE GB GH GH GE IE IT FE LS LU MC MW MZ
         NL GA PT SE SE SE SE NO TRITO UG UM
W: AR AG AL AM AM AM AN BA BA BA BA BA BY BY CA CHICU CO CR CU CZ EK DM
            DI EE ES FI GB GD GE GH GM HE HY ID IL IN 13 SP KE KG KP KE KZ LC
            LK LE LT LU LV MA ND MG ME NU MW ME NU NO NO PL PT RO EU SE SE
            S; SI OK SL TJ TM TH TT TO TA UG US TO VILYU ZA ZW
     AU 2001065863 A 20011030 (200219)
                                                       301N031-31
    THE 10018941 AT DE 2000-10018941 .0000417; WO 2001079315 AT WO 2001-EP3934
     20010406; AU 2001060863 A AU 2001-6506, 2001-406
FUT AU 2001065863 A Based on Wo 200179815
PFAI DE 2000-10018941 20000417; DE 2000-10018940 20000417
TC
     IOM 401N001-05; G01N021-31
     103 - 3017603-36; GU1M001-54
    DE 10018941 A UPAB: 20020321
AR
     MOVELTY - Multiple filter photometer comprises three wide band detectors
     (b) for receiving the transmitted or remitted radiation. The detectors
     mave filters (4) having different spectral binary transmissions of
     approximately u and 1.
          USE - Used for determining small concentration changes in a multiple
     ecomponent substance mixture, e.g. the change of blood glucose in a living
     organism.
```

ADVANTAGE - The concentration changes can be exactly determined. DESCRIPTION OF DEAWING() - The drawing shows a schematic view of the photometer. thermal radiator 1 : ilters 4 wide band detectors ? Lwg.1/8 F.3 TRI EPI FAAB; GI ΜC CPI: J04-B01; J04-C04 EPI: 303-A02A; S0:-E04A; C03-E04B; S03-E04B1A TECH UPTM: 10011211 TECHNOLOGY FOCUS - INCTEUMENTATION AND FESTING - Preferred Arrangement: The number of detectors used correspond to the number of relevant different parameters of the system. The wide band filters are selected so that at least one of the filters lies on a known absorption band of the qual substance. The transmitting spectral regions of the filter are releated so that the Handes in the detector signals caused by substance concentrations and surroundings parameters are maximized. A thermal radiator (1) is used as a light source. L110 ANSWER 4 OF 20 WPIM (C) 2000 THOMSON DERWENT 1001-443245 [48] WPIM DHN 112001-327860 DNC CLU01-134238 Near infra-red spectroscopy online process comprises analyzing constituent figuids against reference spectra data bank of binary mixtures of possible selutions. JU14 803 T01 EK™. BORN, J; FRICKEL, H; ITZEL, H I::(MERE) MERCK PATENT (MBH F/Λ CYC DE 19963561 Al 20010705 (200148)* F'I - 13p - Gő1H021-35 WO 0001048458 A1 00010705 (200148) DE G01N021-31 EW: AT BE OH OY DE DY ES PI FE GB GE IE IT LU MO NL PT SE TE. W: JE US ALT TE 19003861 A1 DE 1900-19868161 19991223; WO 200104:453 A1 WO 2000-EP12199 20001205 PEAT 1E 1999-19963561 19991213 10M G01N021-31; G61N0.1-25 IC IOS G06F017-40 ΑB DE 19963561 A UPAB: 10010919 MCVELTY - A near infra-red spectroscopy chline process analyses the constituent parts of liquid mixtures using a calibration data bank holding the reference spectra of only bunary mixtures of all the possible colutions on presentinea quantitative steps. PETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for an apparatus for the above process. Preferred Features: The pre-defined steps are at irregular intervals of 1-10%. The calculation of spectra consisting of three or four components is effected by a linear combination of the data bank's two-component spectra, taking their respective proportions into account. Evaluation is effected by direct comparison of the compressed full spectral data. Both the spectra measured and the reference spectra are characterized solely by an index number used in consumption with a data reduction module. After data compression and spectral data point scale change and respective wave count, the sum product for each spectrum is redistered, and the differences between the product sums is used for rapid identification of the match between the values measured and the data bank spectra. During data compression, the respective gradient for measured and

USE - The process analyzes the constituent fluid substances in chemical liquid effluent arising from chemical batch production.

and ascending spectral data points and string of numbers.

reference values is coded as a series of numbers, by adding the descending

ADVANTAGE - The process takes less than 30 seconds and is external to and does not interrupt a continuing main process. The process also provides a quantitative indication of water content. Dwg.0.3 CPI EPI

FΑ 44.55

FS

MC CF1: JC4-BolA

EFI: 003-E04A5B; 303-E04A5L; 803-E04U; F01-003B; T01-002; F11-E010; T01-E02A; T01-E02B; T01-H00358E; T01-J04B1; T01-J07A

L110 ANSWER 8 OF 20 WPIN OUR 2002 THOMSON DEFWENT

ΑH 2000-443019 [39] WPIM

DMC 02000-134904 DIN 112000-330448

ΙI Amount of tertilizer to be applied to growing grain crop, its yield and grain quality are ralculated using formula.

EC. 007 P11 P15 803

IIIHOMAKA, Y; MARIYAMA, H; MAKAMURA, N; SATAKE, C

(CATA) ŠATAKĖ CORP; (CATA) GATAKĖ ENG CO LTD; (SATA) JATAKĖ SEISAKUSHO KK FΆ CTC

CA 3250779 A1 [50000369 (L00039)* EN A 1:00.1-00 F. I. $A \cong 1 \oplus \emptyset \cup 1 = \emptyset \oplus$ A 20000316 (200039) AU 9944890 rn: 1251737 A 10000503 (100039) A(0.100, 1-0.0)A-:1-31-7-10 JP 2000300077 A 20001031 (200053) A010021-00 KR 2000023903 A 200004.5 (200107)

ra 1180779 al ca 1999-1.80779 19990826; aŭ 2944890 a aŭ 1999-44890 ADT. 19990401; cn 1251737 A cn 1999-118565 19990303; JP 31 C301077 A CP 1999-154866 19990002; KR 200002/903 A KR 1999-37521 19990903

PRAI JF 1949-154866 | 19990002; JF 1998-254717 | 1998-30009; JF 1999-40080 1.84900.16

1CM A01C021-00; A010007-00

IdS A01G009-00; A(1G010-00; G01N021-31; G06F019-00

ΑВ

MOVELYLY - The amount of fertilizer to be applied to a growing grain crop is raigulated using a formula including information relating the leaf blade to specific periods in the life of the crop, festilizer application related to these periods and information about grain quality. The leaf blace information and target yield of the crop may be entered unto the formula.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the reliewing:

(1) estimating the yield of a grain crop using a formula using the same information as the movel formula;

(i) apparatus for determining the amount of tertilizer to be applied, encep yield or ercy quality, having a memory storing the novel formula, a user data Signit, ar existimatic calculator and an employ display showing the amount of fertilizer to be applied;

(3) apparatus for providing production intohnation of grains, comprising a memory for storing a quality related formula obtained by unalyzing growth information including leaf blade firmation related to specific periods of the crops, fertilizer application information, and quality information of the grains after growth, an input section, an arithmetic section and a display section; and

(4) apparatus for providing production information of grains, comprising a memory storing a yield related formula obtained by analyzing enowth information, and yield information of the grains after growth, an input section, an arithmetic section, and a display section.

USE - The formula can be used to calculate the amount of fertilizer to be applied to a crup, it: yield, and the grain quality of the crop

ADVANTAGE - The grain yield and quality can be estimated accurately before harvest.

.wq. _ , 7

CPI EPI GMPI FS

```
FΑ
     AB; GI; DON
MC
     CPI: C05-A01B; C06-D18; C11-C07B; C12-K04E
     EPI: 30?-E04A5
TECH
                     UPTX: U0070818
     TECHNOLOGY FOCUS - AGRICULTURE - Freferred Features: The leaf blade
     information is the content of hitrogen and chlorophyll, and leaf color.
     These are obtained by spectral analysis. The formula may also use
     information or soil quality.
L110 ANNWER + OF 20 WPIN (C) LOGS THOMSON DEFWENT
    1 * *9-011900 [53] WPIN
AH
ENN N1090-450910
TΙ
    directifier rempensation of Rackground absorption in an atom absorption
     spectioneter.
D:C
     803 TOT UCL VOL
IN BARRWINKEL, W; EICHARDT, E
    (ANAL-M) AMALYTIK JEMA GMBH AMALYCEMMESO(ERAETE
P7.
CTC
                                              15p H01F007-06
    -DE 19816042 - Al 19891014 (199953)*-
Ρï
ADT DE 19816042 AT DE 1998-1981604. 19989409
FFAI LE 1998-19816042 19980403
     IDM H018:07-06
I C
     103 G01J003-42; G01N021-31; G06F007-00
    DE 10816042 A UPAB: 10991215
AL
     MOVELTY - The directit compensates the background absorption using the
     resunctive field of helectromagnet for producing the Zeemann effect in an
     atom abscription spectrometer. The circuit has a parallel circuit of two
     i. r. vortage sources (E1) connected in series and two switches (S1, S2)
     commetted in series with the magnet coil (L) in series with a current
     measurement device (Frg. c. nrected between their junctions.
          DETAILED (BECRIPTION - An INTERENDENT CLAIM is also included for a
     mother of controlling an electric magnet for producing the Zeemann effect in
     an atom absorption spectrometer.
          UCE - For compensation of background absorption in an atom absorption
     spectioneter.
          ALMANTAGE - The woltage supply enables the coil current to follow any
     positive or negative domand value at maximum rate.
          DESCRIPTION OF DRAWING(S) - The righte shows a simplified dircuit
     diagram.
     switches 31, 33
          veitage source El
     Coll L
          current measurement device En.
     I was a C
4(3)
     Direct
\Xi \Lambda
     AB; GI
     EPI: MURIADOE; SOB-EDMAS; TOI-E; MAI-BOIE; MOZ-EDAX
1110 ANDWER 7 OF 20 WRIM (C) 1707 THOMSON DEFWENT
    1999-50%351 [4L] WEIN
M1999-378%53 DNC 01999-148426
AH
DHN
    111999-376853
     on-line measurement of process stream of sugar beet, sugar cane, silage,
Τi
     grain, fruit, regetables, particle board or paper.
\mathbb{D}^{n}\mathbb{C}
     D12 D14 D17 F09 J04 S03
     ATHERTON, P G; BERDING, N; BROTHERTON, G A; GRIMLEY, S C; LETHBRIDGE, P J;
111
     MACKINTOSE, D. L; STAUNTION, S. P; STAUNTOM, S. P.
     (DUGA-N) BUREAU SUGAR EXPERIMENT STATIONS; (SUGA-N) SUGAR NORTH LID
FA
    34
CYC
     We) 9934195 A1 19990705 (199042) * EN = 34p =
                                                     G01N021-31
F'I
        RW: AT BE CH CY DE DE EA ES FI FR OB OH OM OR IE IT EE LS LU MC MW NL
            \odot A PT 3D SE 32 TG 2W
         W: AL AM AT AU AL BA BB BG BR BY TA THICH ON OU OZ DE DK EE ES FI GB GD
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GE GH GM HE HU ID IL IS UP KE NG NP KE KE LC LK LE LS LT LU LV MD

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MG MK MN MW MM NO NZ PL PT RO EU SP SE SG SI SK SL TJ TM TR TT UA
            UG US US VN YU SW
                                              - 1r
     SA 981178? A 19990831 (199942)
                                                     G01N0:0-00
                   A 19990713 (199911)
     AU 9913176
                                                     G01NO. 1-31
     BF 9814406
                  A 20001-10 (200015)
                                                     G 111021-31
                                                                      < - -
                 B 20001150 (200101)
     A0 717034
                                                     G_{i}^{*} 1 No. 1 \pm 3 \pm 1
ADT We 993419: A1 We 1998-A1961 1 09:1117; ZA 9011783 A SA 1995-11753 19981222;
     AU #9121:6 A AU 1999-121:6 19981117; ER 3614416 A BR 1998-14406 19981117,
     We 199: -AT951 199:1117; AU 7: 7074 E AU 13 ee-12186 1 ce 1117
FLT AU 99121a6 A Based on WG 99341b;; ER 9814406 A Based on WG 9934198; AU
     7.7034 B Previous Publ. AU 391.106, Based on WO 9364.93
PRAI AU 1997-1155 19971223
    ICM G01N000-00; G01N021-31
IC
     Ins g015000-00; g017000-00; g017021-13; G06F000-00; G06F000-00
    WO 3934193 A UPAB: 19931014
ΑF
    MCVELTY - On-line measurement of a process stream roads the infrared
     reflectance spectrum of the stream and processes it using a reference
     calibration equation.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the
     following:
          (1) A system having a scanning head with a light source and collector
     or reflected light, a hear intrared spectrophotometer with a monochromator
     for resolving the reflected light into a discrete wavelength, a database
     storing the equation and a computer.
          (in A method of ch-line measurement where the equation statistically
     validates the reflectance spectrum.
          (3) As (a) or (b) where the process stream is sugar case.
          Preferred Features: The parameter is tiber content, juice brix, juice
     polarisation, commercial can sugar, quality parameters, inorganic elements
     or process parameters. Where the material is sugar case it may be at any
     stage from prepared came to crystalline sugar. The spectrophotometer is
     insulated from temperature and vibration by connecting it to the scanning
     head by a fiber optic cable and the head has a vibration damping mounting.
     The system has several spectrophotometers, one of which acts as a standard for the system. The spectrum is 400\,\pm\,250^\circ nano m.
          USE - (all planed) Proceeding sugar beet, sugar came, silage, grain,
     fruit, vegetables, particle board or paper
          ADVANTAGE - intrared spectrum measurements can be made on-line.
     Owg.076
FS
     CFI EFI
FA
     AP
     CFI: DCS-K04; D06-C; EDS-A04C; FDS-A07; JTI-C
ΜC
     RFI: SUS-E04AS
Lilo Spank a object ment to the medway despend
     1994-132789 (21) WRIM
M1994-184374 DNO 01098-072740
A::
DNN N1994-184394
     Projection monitoring using a hear intrared spectrometer - comprises
ΤI
     comparing two groups of spectra using algorithm to cuentify changes in
     product quality.
     H05 J04 S03
EIC!
III
     JABY, C; SABY, C A
     (ERAP) ELF ANTAR FRANCE; 'ERAP' MUF ANTAR MEANCE WA
F'A
CYC
ΡI
                   30111-11-35
         R: AT BE OH DE DE RS FI FR GB GR IE IT LI LU MO NU PT SE
     FF. 1754899 A1 19980424 (199813)
                                                     30111. 1-35
                                              , F,
     CA 2217198
                                                                       < - -
                   A 19980423 (199836)
                                                      60110011-25
                  A .doi:00104 (.do:0005)
     UC 6012019
                                                      G01N U1-31
    EP 333677 A1 EP 1947-402493 19971021; FR 2754899 A1 FR 1996-12917
     14961023; CA 2217124 A CA 1937-2217108 19371022; US +012019 A US
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1897-956436 19971025 PRAI FE 1996-12917 19961023

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{\rm IC}
     ICM G01N021-25; G01N021-31; G01N021-35
     ICJ
         G01D018-00; G01N033-28; G01N037-00;
          G06F017-00; G06F017-10
           938077 A UPAB: 19380505
ΑE
     EΡ
     Following and surveying the function of a unit fabricating a product
     and/or a near infrared spectrometer fed by the product, where the
     spectrometer delivers spectra comprising a series of absorbence values for
     different wavelengths, comprises: (i) recording the spectra from the near
     infrared opentrometer periodically in the form of numerical data; and (ii)
     trunsforming the data from each spectrum mathematically to obtain
     *ransformed spectra. In addition the following stages are used: (i) a
     sories of working spectra are made from the obtained spectra by choosing
     the wavelengths in each transformed spectrum by a method of selection;
     (ii) a first set of 20-56 comp outive spectra are selected from the
     working spectra and a second set of spectra the same size is selected so
     they are consecutive and out of phase to the first set; and (iii) at least
     one quality priterion is calculated to compare the two sets of spectra and
     the evolution of this oriterion is followed over time.
          USE - Used for monitoring industrial chemicals, petrochemicals,
     pharmaceuticals, foodstuffs etc...
          ADVANTAGE - The process is easy to use and identifies changes in
     product due to dysiunchions.
     Lwg.1/3
    CFI EFI
FS
     AF: GI
FA
MC
     CFI: H05-H; J04-CUS
     EFI: 003-A02; 003-E04A0; 00:-E04A5B; 003-E14A; 003-E14A1; 003-E14F
L110 ANGWER 9 OF 20 WPIX (C) 2001 THOMSON DERMENT
     1947-297819 [27] WPIK
AΩ
DNN 111337-246132
    Instrument for optical measurement of living body - has several light
I I
     modules emitting light into several positions of body through optical
     typres, with beams of light transmitted to surface of body picked up at
     several positions by photodetectors.
DC
     F-1 803 205 T01
     ROICUMI, E; MAKI, A; YAMASHITA, Y
TIL
PA.
     (HITA: HITACHE LTD; (FOIZ-I: ROIZUME H; MAKE-I) MAKE A; (YAMA-I)
     TAMASHITA Y
CYC
    ۲,
                   A1 19976529 (189729)* JA 1 (p) A618/08-14
PΙ
    No. 0718751
         W: CA DE GB U.1
                  A 199705U7 (199731)
                                               1.4
     JP 49138825
                                                      A61P005-14
                   A
                       19970610 (199733)
                                                      2615 005-14
     JF 09143894
                                                1-1
                       19970610 (199737)
     TF 09149909
                   A
                                                ** 1
                                                      A618010-00
                   A 19971008 (199748)
                                              \mathbf{T}
     GH 7311884
                                                      G01N021-1
                                                                       ·# --
                   T 10071011 (199804)
B 000000322 (200018)
                                                       \Lambda(i,1\otimes i)((i\otimes -i)):
     DE 19681107
                                                      G(\{1N\}) \subseteq I = I^{-1}
     GB 12311354
     TO (240309 B1 20010519 (200133)
                                                       7.6.1\,\mathrm{E}\,\mathrm{M}^{-1}\,\mathrm{G} = 3.5
                                                      7911B0011=00
     US 2001018554 Al 200105 0 (200151)
                                                      A618965-14
     CA 121070: C L.U11009 (LUU.03) EN
ADT WO 9718755 AT WO 1996-JPE365 19961115; JP 09185825 A JP 1995-299542
     14951117; JP 09143694 A JP 1995-314195 1 001201; JP 09149007 A JP
     1995-311993 19951150; GB 2311894 A WO 1996-725365 19961115, GB 1997-13904
     19970019; DB 19681197 T OE 1996-196:1107 19961115, WO 1996-798665
     19961115; GB 2511454 B WO 1996-CP3565 19961115, GB 1997-19094 19970619; US
     6240309 B1 CIP of US 1995-539871 19951006, WO 1996-JP3365 19961115, US
     1990-875981 19970929; UN 1001018554 AT CIP of US 1996-539 71 19951006,
     dont of US 1997-375031 19970429, US 2001-40409 20010507; CA 2210703 C CA
     1996-2210003 19961115, WO 1996-3P3365 19961115
    GB 0311854 A Based on Wo 9718755; DE 196-1107 T Based on WO 9718755; GB
     241854 B Based on WO 9718755; US 624030+ B1 CIP of US 5803 409, Based on
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Wo 9718755; US 2001018554 A1 GIP of US 5003909, Comb of US 6240309; CA

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2010703 C Based on Wo 9719755
PRAI JP 1995-714195 19951201; JP 1995-299542 19951117; JP 1995-211993
     19951130
EEP AU 5416934; DE 4314685; DE 431523; DE 4393333; DE 136392; DE 44693; DE
     49793; EP 187530; EP 319158; EP 614645; EP 619055; FI 961673; JP 1262839;
     UF 3.41414; CP 3605322; 3P 4168144; JP 5103186; CP 5118438; GP 5261110; JP
     8/115231; JP 61502660; JP 6181850; JP 62251610; JP 6245978; JP 630C5234;
     urp 03305a45; dp 7103a64; dp 7117595; dp 7120384; dp 7204168; dp 3233727;
     JF +019408; NO 351730; US 4576173; US 4800380; US 5408630; WO 8600514; WO
     8912223; WO 9405209; WO 9410301
     icm AsiBoc1-os; AsiBous-48; AsiBCDS-14; AsiBo06- 0; AsiBc10-DD;
ΙĊ
          G01N021-17
         A61E001-00; A61E005-00; A61E006-03; F61E023-36; G01N021-27;
     103
          G01N021-31; G01N033-49; G06F003-00
          9718755 A UPAB: 19970731
A.E.
     The optical measurement instrument has a light source (I) which includes
     several light modules (92(1) to 2(10)) which emit intensity-modulated
     beams of light at different frequencies through optical fibers (f-1 to
     w-10) so that they can be introduced into the living body (3) at several
     points. The beams of light passing through the body are picked up on the
     surface of the living body (3) and quided to photodetectors (11-1 to
     11-25) through optical fibers (10-1 to 10-25).
          The signals from the photouetector (11-1 to 11-105) are imputted to a
     look-in amplifier module (11), where the intensity of the return beam
     detected by each of the photodetectors and having the same modulation
     frequency as that of its corresponding input beam is selectively measured.
     The intensities of beams of light picked up at several positions are
     richessed by a data processor (16).
          ADVANTAGE - Internal information of loving body for several pick-up
     resitions can be obtained without prosstack.
     Ewg. 047:7
F.
     EFI GMFT
FÆ.
     AB; GI
     EPI: $03-E04C3; $05-D0.X; T01-J16A
MC
L110 ANSWER 10 OF 20 WEIN (C) 2002 THOMSON DEFWENT
P.::
    1987-214928 [20] WPIM
ENN H1997-177184
     Controlling working of analyser and fabrication unit for control
ΤI
     laboratories e.g. petroleum - ising multi variate dalibration of master
     analyser, periodic standardisation of slave analysers and calibration
     transfers between analysers.
[:t]
     S03
I::
     \text{EI} \cap \text{UE}, \text{F}; \text{SABY}, \text{C} A
     SEMAPO ELF ANTAR PRANCE: (BRAD RIF ANTAR BRANCE 'A
FΑ
\mathbb{C} \mathbb{T} \mathbb{C}
         0351. AL 19300410 (1937.171 FE
H: BE CH DE DE ES GE IT LI NL SE
ΕI
     EF 753511
                                                 - 7p - G 112 h1-. 7
                  A1 14470418 (134728)
                                                        G. 11 11 - C
     FF. L 139923
                   A 19970417 (199733)
A 19970711 (194735)
     CA 2137043
                                                        G:110(18-00
                                                        GHIMME - M
     JE 0017±75€
                                                 1171
                  As 199710..2 (199814)
A 20000181 (200015)
A 19001003 (200050)
                                                        G 111 (111-17
     EP /68801
                                                        G01H021-00
     II. 119427
     US 6118544
                                                        306F-119- ()
ADT EF 768522 A2 EP 1996-402187 19961015; FR 1739928 A1 FR 1995-12087
     1496-1016; CA 1187945 A CA 1496-1187945 19461015; JP 08178756 A JP
     1346-304853 19961016; EP 768522 A3 EP 1996-402187 14961015; IL 119427 A IL
     1996-119427 19961015; US 6129544 A US 1996-732117 19961015
                      13351016
FEAI FE 1995-1.037
REP In-Sk.Pub; 1.5ml.Ref; US 4806644; US 5243546; US 5459677
     I:M 3010013-00; G01N021-00; G:01N035-00; G06F019-00
     ICS - 5010021-00; G01N021-31; G01N021-55; G06F015-18;
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G06F017-10

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ICA G01N001-27
AΕ
    EP 768502 A UPAB: 19976516
     The method of following and controlling the operation of a slave analyser
     linked to a fabrication unit area an operation of multivariate calibration
     of the master analyser and periodic standardisation operations of signals
     delivered by the slave analyser fed by products of standardisation.
          Calibration transfer operations letween the master and slave
     analysers are undertaken including the calculation of parameters
     associated with a dalibration transfor algorithm and the choice of a
     control indicator. The evolution of the indicator is reviewed at the start
     of each periodic standardisation operation and used to check the correct
     working of the slave analyser and fabrication unit.
          USE/ADVANTAGE - Also for research laboratories and manufacturing
     units in themical, pharmacoutical, occuratic, food and agricultural
     industries. Method enables causes of drift and malfunction to be
     identified.
     fiwa. 0.0
F.,
    E.E.I
FA
    AE
     EPI: :003-E04Al; S03-E04T
MC
L110 ANSWER 11 OF 20 WPIK (C) 1002 THOMSON DEEMENT
     1994-351270 [44]
                       WPIK
DMN 111994-275640
ΙT
    Instantaneous spectroscopic analysis system for fluid - uses correlation
     of spectrum produced with those held in memory to determine relative
     concentration of various products in cluid sample.
DC.
     803 T01
III
     FACHINGER, C: MARTIN-BOUYER, M: NAFFRECHOUK, E: SUPTIL, J
     (UYSA-N) UNIV SAVOIE
EV
C':C
E. I
     FF 1704650
                Al 13941164 (199444)*
                                              1 350
                                                     -G01N021-27
     WO 94058-7 A1 19341110 (193444) EF
                                                     \oplus \{1,r\} \mid \exists - \exists \exists
     EF 647316
                 A1 19950412 (199519) FY
                                                     G017013-23
     US 5528363
                                                     G01J063-18
                 A 19960618 (199630)
                                               ńρ
ALT FF 0704650 A1 FR 1993-5204 1993-3204; WO 9425837 A1 WO 1994-FR408 19940427;
     EP 647310 AL EP 1994-914451 19940427, WO 1994-PF478 19940427; US 5528363 A
     WG 1994-FF478 19940427, US 1994-360744 19941222
FIG. EP 647316 Al Based on WO 9425837; US 1528363 A Based on WO 9425837
FFAI FF 1993-5364
                     13330437
    -06Jnl.Ref; EP 368500; EP 309057; EP 510322; GB 2217006; WO 9013810
     ICM G01J003-18; G01J003-28; G01N021-27
IC
     ICS G01J003-50; G01N021-31; G06F015-20
         02704650 A UPAB: 19941025
AВ
    E'F
     The detection and identification system operates by illumination of the
     medium being examined by a memochromatic or polychromatic light wave (A),
     and processing of the resulting emission, abstrption or reflection
     spectrum. This is achieved with a plane field monochromator (5) and a
     transformation circuit (6: producing analogue or digital signals.
          The circuit is ocupled to a central processing unit (7) analysing the
     spectrum produced for comparison and correlation with spectra stored in
     memory. As a result of this comparison, the concentration of various
     physics-onemical products within the redium may be determined.
          ADVANTAGE - System provides instantaneous analysis of composition of
     fluid.
     Dwg.1/1
     EPI
F.5
FΆ
     AB; GI
     EPI: NOS-E04A1; T01-J05A
MC
ABEQ TS - 9528 WEE A UPAB: 1996 (731
     A compact portable device capable of operating in a hostile environment to
     carry out qualitative and quantitative identification of one or of a
     plurality of physicochemical entities contained in a sample capable of
```

producing a spectra under expitation by electromagnetic waves that includes:

a portable compact casing that shields equipment housed therein from magnetic fields, electric fields, and external pressure variations;

a polychromator having an optical path within said dasing to which emissions, absorption or reflective spectrum of a liquid, solid or gas sample are transmitted and analysed, having said spectrum being decomposed into a sequence of signals having discrete variation in vavelength;

detecting means for detecting said discrete wavelength signals positioned on the optical path of the polychromator;

conversion direct means operpled to waild detecting means for polyerting said discrete wavelength signals into electrical signals; and,

probessing means having a plurality of standard spectra represented of anown entities in memory for analyzing the spectra embodied in said electric signals and to deportelate and compare the spectra with that stored in memory and to determine the nature and concentration in said sample.

```
Dwq. 1/1
L110 ANOWER 12 OF L0 WPIM (C. 2002 THOMSON DERWENT
     1904-279897 [34] WPIM
DHN N1994-310459
                           PRIC (11994-12779)
TI
     Optical measuring unit for modifications in reactive substance in
     transparent cell - passes light through cell and then through filters to
     brightness measuring elements.
DC
     JG4 S03
III
     DELIGNIERES, E; DURAND, C
     (INSF) INST FRANCAIS DU PÉTROLE
FA
CYC
                     Al 1994051a (199434):
     WG -4114543
                                                    1.1
                                                            G.1NOL1-*1
ΕI
         RW: AT RE CH DE DK ES FR GB GR IE IT LU MC NL PT SE
          W: CA JP TO
     FR 2701318 A1 19940812 (199404)
EB 288127 A1 1994012 (19850)
                                                             G[MEL1-i1]
                                                     1 (Eq.)
                     - Al (1395011) (139505) - 89
                                                             G11111. 1-51
          F: BE DE GE IT NL SE
      JE 98501394 - W 19960217 (199643)
                                                      1 340
                                                             GP1100, 1-09
     US 8650220 A 19971021 (199748)
EE 639127 B1 19980916 (199841) FP
                                                      10p
                                                             G111011-13
                                                             GP10001-31
          F: FF DE GP IT NL SE
DE 69413331 E 1998102. (199849) GC10021-31 R-ADT WC 9418543 A1 WO 1994-FRIGO 19940191; EN 1701313 A1 EN 1998-1513 19930209;
     ER 635127 A1 ER 1994-905765 19940171, WO 1994-FR120 19940131; JP 68501394
     W of 1994-51771 19340131, Wo 1904-FR12, 19941131; US 5680220 A WO 1994-FR130 19940131, US 1995-30778, 19940137; EF 635107 B1 EF 1994-905765 19940131, WO 1994-FR120 19940131; DE 69413331 E DF 1944-613331 19940131,
EF 1994-905765 19940131, WO 1994-FR120 19940151
FDT EF 035117 Al Ruwed on WO 9418141; OP 03101134 W Based on WO 3418543; US
      56%0226 A Basou on WO 9418543; EF 03512 Bl Based on WO 9418343; DE
      69413331 E Based on EP 035127, Bused on WO 3418343
FRAI FR 1993-1513 19930209
     US 3902776; UJ 3925540
F.E.P
      IC
      ICN G017003-46; G01N021-27; G06F015-46
           9418543 A UPAB: 19941013
АБ
     W()
      The unit consists of at least one light source (1) with two optical
      branches (11,13) which allow the incident light to be passed selectively
```

The unit consists of at least one light source (1) with two optical branches (11,12) which allow the incident light to be passed selectively through the cell (3), an optical system (7,8) to direct the emergent light rays through selective optical filters (51, 52, 53) of different wavelengths, and measuring elements (51, 52, 53) for the intensity of light passing through the filters.

The emergent light bays from the cell and a neutral filter (ξ) can be passed selectively through the filters by means of two shutters (01,02) and a switching system (M,I). The unit also incorporates a control

assembly (9) with a command processor (10), a signal acquisition unit (11), and an interface assembly (12).

USE/ADVANTAGE - Appts, can be used for determining the pH of a substance. More precise and reliable results are obtd. Ewa.1/6

FSCFI EFT

AB; GI FF.

Mc" CPI: U04-C

EFI: 307-E94A5

AEEQ US 5680220 A UPAB: 19971209

A devise for optically measuring modifications in a reacting substance contained in a transparent cell, comprising

a single light course provided with an electric supply voltage and a specified light spectrum, a first optical circuit, a first optical shutter arranged in said first optical curruit, a second optical circuit, a second aptical shutter arranged in said second optical directit, an optical diverter in said optical circuits for diverting incident light from the single light source through the Lell and through a reference medium to an optical mode, an optical separator for directing light from the optical node to three other optical circuits, a set of three selective filters arranged respectively in the three other optical circuits, a first selective filter from the set being centred on a first wavelength corresponding to an isobestic point of the reacting substance, a second selective filter from the set being centred on a wavelength in a part of the light spectrum where the reacting substance is the most sensitive and a third selective filter from the set being centred on another part of the light spectrum where the reacting substance is the least densitive, a measuring means for respectively measuring the Light emanating from the three other optical circuits, including a set of three detectors for respectively detecting light passed by each of the set of three filters and producing cutput signals representing the detected light, an electric power supply for providing the electrical supply voltage, a controller and an electric switching means controlled by the controller for connecting intermittently the three detectors to the controller, for connecting the single light sturbe to the power supply, and for selectively switching the optical shutters. Involution

L110 ANSWER 13 OF 20 WPIM (C) .002 THOMSON DERWENT

AN 1994-201191 [25] WPIN

DIIN N1994-15-247

ΤI Spectrumeter with dynamically coded components - has data carriers, readers, writers and central computer for coded data characterising replaceable components.

DC: 503

III FARR, N; SIMON, A; WEIL, J

P7. PRUKHM) BRUKER ANALYTI. OHE MESUMECHNIK; (BRUKHN) BRUKER ANALYTISCHE MESSTECHNIK GMBH

CYC

DE 4041305 A1 19940616 (1994.5)* G01J003-02 310 PΙ C2 19950126 (199509) A 19960917 (199648) DE 4241 305 7:5 -G01J083-02 H11J00E-08 7 p 03 9557544

DE 4041905 A1 DE 1990-4041905 10921211; DE 4241905 C2 DE 1990-4241905 19921211; US 5557544 A US 1993-164390 19931209 PRAI DE 1992-4841305 19981211

ICM GC13033-02; H017005-02 ICB G01N021-31; G01N024-08; G01N024-13; G01N027-62; GC2B027-00; G06F013-00

Ais -4.41.05 A UPAB: 13.40810

The spectrometer has a bentral computer, a radiation source, detector, mean divider, filter and external measurement probe. The replaceable components each have a readable data carrier (7) with coded data contg. the parameters characterizing each component. The data carrier is I ship, esp. an EPROM or a flash-FOM.

The data carrier can be written into and contains variable, time dependent data concerning the prewritten and current characteristics of each replaceable component, e.g. operating duration, wear parameters or calibration curves. The central computer decodes the data and derives deviations.

UUE/ADVANTAGE - Exp. for infrared analytic spectrometer, bit also MME, EUE, ICR, or mass spectrometer. Dynamically coded components can be used with great flexibility at other points in dame spectrometer or in their spectrometers.

Dwg.172

FS EPI

FA AB; GI

MC EFI: 003-A02B; S03-E04A5; S03-E07; S03-E10A

ABEQ DE 4241005 C UPAB: 19950301

An analytical, esp. IE, spectrometer has a CPU (3), fixed components and replaceable components (5) including a readable data carrier (7) with coded data on component parameters. These data can be read and transmitted (8a) to the CPU which has a decoding and decision-making programme.

one or more devices input variable time-dependent data into the data carrier on the actual condition of the replaceable components. The CPU has a programme to control the data writing device and to automatically match the data on changed component parameters on its data carrier.

ALVANTAGE - Greater component flexibility at different positions in the same or other spectrometers.

Dwg.27.

ABEO US 5557544 A CEAB: 19961025

An FTIR spectrometer comprising:

an interchangeable optical component;

readable data medium means integral with the interchangeable optical component and adapted for storing encoded data concerning at least one of a mistory and a changeable current property of the interchangeable optical component and adapted for storing non-changeable encoded data;

read/write means connected to the data medium means for reading encoded data from and for writing encoded data to the data medium means;

a central computer adapted to decode the encoded data and to process decisions on the basis of decided data and adapted to process the encoded data for controlling the read/write means to automatically update changed parameters of the interchangeable optical component;

interface means connected between the read write means and the sentral computer for transferring the encoded data to and from the sentral computer; and

sensor means, communicating with the read/write means, for detecting changes in the encoded data and for generating data signals in response to the detected changes in the encoded data bwg.1/.

L110 ANSWER 14 OF 20 WPIK (C) 2002 TROMSON DERWENT

AN 1993-218849 [32] WEIM

DHN 11993-199136

TI Instrument for non-destructive measurement of material properties - uses data obtd. from material by sections of electromagnetic spectrum to determine material properties by data-fusion analysis.

DC 803 T01 E11

IN ESCHERGAR, E P

PA (SGII-H) SGI INT

CYC L8

PI WO 9515470 A1 19930808 (199332)* EN 82p G36F115-46 <--

 W: AU BR CA JP KE N2 PL EO EU UA

 AU 9335940
 A 19930001 (199310)
 \$16F015-46

 US 5291412
 A 19940301 (199400)
 13p
 \$C1N021-47

 JP 07507133
 W 19950803 (199539)
 10p
 \$301N022-00

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ADT WO P315470 A1 WO 1993-US764 19930126; AU 9335940 A AU 1993-35940 19930126;
    US 5291432 A US 1993-826780 109.0128; JP 07507133 W JP 1993-513396
     1 +3 40126, WO 1393-T3794 13 430126
    AT 93 0 940 A Based on Wo 9:15470; JP G7507133 W Based on WO 9:15470
FFAI US 1992-8.6780 19920123
    UD 04/57201; US 4616317; UN 4-55759; US 5127268
F.E.P
     ICM G01N021-47; G01N00L-00; G06F015-46
IC.
     IC3 G01N021-31; G41N023-22
         9315470 A UPAB: 13940510
A.E
    Samples (14) are carried by conveyor belt (16) past an array of sensors
     (20), each of which stimulates electromagnetic spectrum from the samples
    and detects that radiation. Each sensor has an associated microprocessor
    controller (22) which feeds detected data to a data processing computer
     (11). The data processor merges the data from the sensors by matrix
    techniques and passes the results to the user via a communication path
     (m.6). The user can also supply instructions and additional data e.g. from
    other forms of measurement to the data processor.
         MCE/ADVANTAGE - Analysis of e.g. soal, food products, wood, dement,
    petrologm. Multiple measurements performed automatically on a simple
    sample eliminate need for multiple samples, increuse accuracy and save
    both time and effort. Properties difficult to measure directly may be
    measured indirectly more easily and the required result obtd. by data
    fusion techniques.
    Dwg.1/4
    Ewg.1/4
F\mathcal{E}
    EFI
F'A.
    AB; GI
    EP1: 803-E04A; 803-E05; 803-E06; 803-E07; 803-E14E1; T01-868A; K11-A09
ΜC
The non-destructive, non-contacting instrument detects electromagnetic
     radiation from the materials apposs a wide range of the electromagnetic
    spectrum, and combines these diverse data to derive the material property
    values desired.
          In particular, the properties detectable through particle magnetic
    resonance, spectroscopy of light in the infrared-visible-ultraviolet
    range, and detection of K-ray and gamma ray radiation may be included in
    the instrument. Sensors detect each wavelength band of electromagnetic
    radiation, and data from all these sensors are merged in a fentral data
    processor to evaluate the material properties of interest.
         USE/ADVANTAGE - Frovides material property measurements quickly and
    automati milly, using single sample of test material. Incorporates data
    fusion to enable information about material to be derived by correlation
    of disparate sensor data, with minimal human intervention required.
    Dwg.1/4
L110 ANSWER 15 OF LO WRIN (C) 1000 THOMSON DERWENT
AN
    1992-116207 [15] WPIM
    Spectroscopic determin, of one constituent in fluid mixt. - such as oil
ΤΙ
    content of wax, suitable for on-line or batch measurement, avoiding need
     for dilution.
DUD
    H02 J01 J04 S03
    CHIMENTI, R J L; HALPERN, G M
IN
    (ESBO) EMMON HES & ENG CO
FF
CYC
F'I
    EF 479472
                  A 14420408 (199218)*
                                             1 15
        F.: DE FR GB IT
    CA 2050108
                 A 19920327 (19923)
                                                     GH1N021-15
                  A3 199.00fd) (199351)
     EF 479475
     T. 53-41125
                  A 1994(4): (1994(5)
                                             100
                                                    3 (6F015-00
                  B1 19950614 (199518) EN
     EP 479471
                                             4p
                                                    G01N021-25
        F.: DE FF GB IT
```

G01N021-35

GJ1ND21-25

DE 69110390 E 19950020 (199504)

CA 20:010a

S 2 010913 (20 157) EN

ADT CA 2050108 A CA 1991-2050108 19910828; EP 479472 A3 EP 1991-308682 19010924; US 5301125 A US 1990-588649 19900926; EP 479472 B1 EP 1991-308682 19910924; DE 69110390 E DE 1991-610390 19910924, EP 1991-308682 19910924; CA 2050108 C CA 1991-050103 19910828

FDT DE 69110390 E Based on EP 479472

PEAI US 1990-588649 19900926

REP No-SR.Pub; DE 3625490; GB 20200009; US 4449810; EP 3625490

AB EP 479472 A UPAB: 19931118

Process is for spectroscopic determination of amt. of one constituent of a fluid mixt. in another constituent or in the fluid mixt. itself, following seph. of the mixt, into its constituents, in which a spectroscopic determination of the amount alpha Nn of the Nth consituent of a fluid mixt. 0 in another constituent n of the mixt. following the sepn. of the mimt. Into M constituents 1, ...M (where M i sup to M) and where, due to imperfect seph., the amt. alpha of constituent M remains present with separated constituent n, comprises (a) determining the absorptivity aN of constituent N at a selected wavelength, or at multiple wavelength, across a selected wavelength range, in which constituent N exhibits light absorption; (b) determining the absorptivity an of the other constituent n with the amt. alpha of constituent N present at the same selected wavelength or by the same multiple wavelengths; and (c) determining the amount alpha of the one constituent Npresent with constituent n from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities an and all whore the absorptivities are expressed solely as the ratio of an/aN.

USE/ADVANTAGE - The method is suitable for measuring the oil content of wax, and is suitable for on-line measurement or batch measurement, as circumstances require, the method avoiding the need for dilution. The use of the method for determining the oil content of wax obtd. By dewaxing of oil boiling in the lubricating oil range is specifically claimed.

5.78

ress

FS OPI EPI

5/0

FA AB; GI

MC CPI: NO5-K; J04-P01A

EPI: S03-E04A5; S03-E04B1A

APEO EP 479472 A UFAB: 19931006

Process is for spectroscopic determination of amt. of one constituent of a fluid mixt. in another constituent or in the fluid mixt. itself, following seph. of the mixt. into its constituents, in which a spectroscopic determination of the amount alpha Nn of the Nth constituent of a fluid mixt. O in another constituent n of the mixt. following the seph. of the mixt. into M constituents 1, ...M where N is sup to M' and where, due to imperfect seph., the amt. alpha of constituent N remains present with separated constituent n, comprises (a) determining the absorptivity aN of constituent N at a selected wavelength, or at multiple wavelength, across a selected wavelength range, in which constituent N exhibits light absorption; (b) determining the absorptivity an of the other constituent n with the amt. alpha of constituent N present at the same selected wavelength or by the same multiple wavelengths; and (c) determining the amount alpha of the one constituent Npresent with constituent n from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities an and aN where the absorptivities are expressed solely as the ratio of an/aN.

USE/ADVANTAGE - The method is suitable for measuring the cil content of wax, and is suitable for on-line measurement or batch measurement, as circumstances require, the method avoiding the need for dilution. The use of the method for determining the bil content of wax obtd. By dewaxing of oil boiling in the lubricating fil range is specifically claimed.

ABEO US 5301125 A UPAB: 19940517

When separating a fluid mixture into fractions, the amount of the Nth constituent remaining in another constituent n is determined by measuring the light absorption of N and n and determining the amount of N in n using an expression in which the absorptions are expressed solely as the ratio and aN. The method is partic, for determination of the entrained oil content of wax resulting from separation of a waxy raffinate into dewaxed hydrocarbon oil boiling in the Loricating oil range, with solvent added to samples of wax and oil fractions before the determination.

ADVANTAGE - Simplifies determin, and can be used on-line or in a batch process. Own.378

ABEQ 50 47.4472 B MMAB: 13050721

A method for the spectroscopic determination of the amount alln of the Nth constituent of a fluid mixture O in another constituent host the mixture tellowing the separation of said mixture into M constituents 1,...,M (where n, N is less than M) and where, due to imperfect separation, said amount all of constituent I remains present with separated constituent n, said method communising the steps of:- it determining the absorptivity aN of separated constituent M at a selected wavelength, or at multiple wavelengths, across a selected wavelength range, in which reactituent N exhibits light absorption; (ii) determining the absorptivity an of said another separated constituent h with said amount aNn of constituent N present at the same selected wavelength or at the same multiple wavelengths; and (iii) determining the amount aNn of said one constituent If present with constituent in from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities on and all where the absorptivities are expressed solely as the ratio andaM. Invo.57%

L110 ANSWER 19 OF 20 WRIN 50 2001 THOMSON DERWENT

AN 1391-312,564 (43 WEIM

DNN N1 991-139399

TI Atomic absorption spectrophotometer, dalibration method - uses comparative technique for determining constituent of sample by measuring and storing several standards nominal values.

DC 391 803 Tel

IN MASSAFT, D; MASSART, E U L

FA (PHIG) PHILIPS GLOEILAMPENFAB NV; (PHIG) PHILIPS ELTRIN & ASSOC IND LTD; (PHIG) KONINK PHILIPS FLECTFONICS NV; (PHIG) PHILIPS ELECTRONICS UK LTD; (PHIG) US PHILIPS COPP

CYC

ΕI EP 453036 A 19911023 (199143)* F: CH LE FF GB LI A 10811010 (100147) A 19911024 (199150) A 19930511 (199310) B 18940418 (199411) GF ...4...11 AU +175075 113 5010778 (30)11018 - [6]3p ATT 646545 (30,110)(1,1-31)UD 5880997 UU 5551997 A 19960 00 (190641) BP 453036 P1 19991201 (200001) EN 19960 *0- (199641) G12E015-E0 10p $\oplus 0.110011 - 11$ F: CH DE FR GB LI DE 69131806 E 20000105 (20000) G011101 1-21 . . . _ _ JP -081270 BL 20000828 (100044) -601101.1-17 Эp

APT ER 453036 A EP 1991-200880 19910415; GB 2243211 A GR 1990-83.2 19900410; UC 5210778 A UC 1991-685.36 19910412; AU 648545 B AU 1991-75073 19910418; UC 5951997 A Cont of UC 1991-85266 19910412, Cont of US 1992-976624 19911116, UC 1994-275099 19940714; EP 453036 B1 EP 1991-200830 19910415; DE 091718006 E DE 1991-001800 19910415, EP 1991-200880 19910415; CP 3081270 BC 5P 1991-415339 199104.0

FPT AT 648545 B Previous Pub.. AT 9175073; US 5552997 A Cont of US 5210778; DE 64151806 E Based on EP 453036; JP 5081270 B2 Previous Publ. JP 04230834

PFAI GB 1999-8922 19900420

REP 4.Jnl.Ref; A3...9150; DE 3406_23; NoSE.Pub

G01N021-31; G01N023-06; G06F015-20 ICM G015018-00; G01N021-27; G01N021-31; G12B013-00 ICS G011023-06; G06F015-20; G06F019-00

ICA G01N023-20; G01N023-22

45 (0)6 A UPAB: 19930928 AB

The method of calibrating an analytical instrument which uses a comparative technique for determining a constituent of a sample involves measuring a characteristic of several standards having different nominal values. The measured characteristic is stored in association with its corresponding numinal values. A best straight line is determined using statistical techniques on the stored values. The quality of the calibration line is determined. When the quality of the calibration line is not acceptable, the slope of the line joining each of the stored measured characteristics and nominal values to the origin is determined. It is also determined whether the slopes have a given order.

The method comprises determining the slope of each stored measured characteristic and nominal value with respect to the first measured characteristic and nominal value. The curvature of the calibration line is determined if the slopes have a given order. If the slopes have not a given order it is indicated that the calibration line does not pass through the crisin.

ADMANTAGE - Enables determination of possible dauses of lack of quality of calibration line, by funding whether or not slopes are random so determining whether the problem is in precision of measurement of standards or quality of standards, or whether the points do not represent straight line through origin, but represent duried line.

1/7

FЗ $\mathbb{E}\mathbb{P}\mathbb{I}$ FAAB; GI

EFI: S02-K09; S03-E04D; T01-J MC5310778 A UPAB: 19931115

The atomic absorption spectrophotometer has advice for measuring the absorbahoo of a number of standards of known concentration (110) and plots the measured absorbance against concentration (III). A straight line is fitted to the protted points (102) and a quality coefft. calculated (103). If the quality obetft, is acceptable (104) the balabration line is used for measurement of samples (105). If not, then the slope of the line joining mach point to the origin is determined and if the slopes are random (197) them a remust regression technique is used to fit the palibration line (10s). If outliers are then detected (11s) it is determined which points are outliers (110) and appropriate action taken, for example to restrict the range if the last point(s) is/are butliers (111).2/7

If the aleged ditermined in easy (199) are not random, than, provided more than four points remain (11%), the slope of each point with respect to the first point is determined (114). If they are again not random (115), then a curved calibration line is diagnosed while if they are random, a straight line not passing through the origin is diagnosed (117). In atomic absorption spectroscopy a straight line not passing through the origin indicates a problem with the plank solution, for example, contamination.

USE - For determining themical properties of sample.

Dwg. 277 5553997 A UPAB: 19961011

A method of calibrating an M-ray spectrometer comprising the steps of

(a) generating x-rays from at least one standard sample,

- (b) measuring intensities of said x-rays for different concentrations of said standard sample,
- (c) forming a representation of intensity versus concentration for each measured value of intensity with said different concentrations,
- (d) determining if a best straight calibration line can be formed from said representation, and if not

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(e) changing instrument parameters and/or sample preparation to
     achieve said best straight calibration line, and
          (t) repeating said steps (a)-(d) to calibrate said x-ray
     spectrometer.
     Dwq.1/7
L110 ANSWER 17 OF 20 WPIM (C) 200. THOMSON DEEMENT
    1989-001+53 [01] WHIK
DNN N1989-001581
    Optical interference, absorption or statter determining appts. - has
TI
    measured values of electromagnetic radiation intensity correlated with
    different sets of known values derived from model.
DC
    S02 S03 T01
III
    EDGAR, R F
    (INFR-N) INFPARED ENG LTD; (INFF) INFRARED IND INC
FΆ
CYC 6
E. I
    GB 2206429
                  A 19890105 (198901) *
                                              . Jp
                  A 19890118 (198903) EN
    EP 299646
        H: BE DE FR GB IT
                  A 19900828 (199037)
     D3 4952061
                  B1 19930505 (199318) EN
                                              1 ép
                                                     G01N021=31
     EP 299646
        R: BE DE FR GB IT
                  G 19930609 (199324)
                                                     G01N001-31
     DE 3880748
ALT GB 0006409 A GB 1987-18608 19870700; EP 289646 A EP 1868-505915 19880629;
    US 4952061 A US 1988-211708 19880627; EP 399646 B1 EP 1988-305915
     19880629; DE 3880748 G DE 1985-388074% 19850629, EF 1 688-305915 19880629
FDT DE 2880748 G Based on EP 209640
PFAI GB 1987-15608
                     19870702
REP Att., 9033; DE 2426598; EP 230305; No-SR. Eub; SU 123 (508; US 4655/67;
    Olumbia. Ref
    G01B011-02; G01N021-31; G06F000-01
IC.
    ICM G01N021-31
     ICS G01B011-02; G01M021-35; G01N021-41; G01M021-47; G01M021-38;
          G06F000-01
         3206419 A UPAB: 19930923
ΑĿ
     Electromagnetic radiation is transmitted through or reflected from a
     sample. The radiation includes at least two spectrally different
     components so taht at least one of the components is subjected to optical
     interference, absorption or scatter. The components are transmitted
    through or reflected from the sample by respectively different amounts.
          The transmittance or reflectance of the sample for each of the
    components is measured to derive respective measured values, before
     correlating by either a zero dependent correlation function (\hat{s}), or a
     residual function (Nres). Known vacues having an otpinum correlatin with
     the measured values are selected, with the selected values representing
     the property, or the identity of the sample which is sensed or to be
     determined.
          ADVANTAGE - Both functions are unaffected by dair factors, thus
     avoiding any need to determine and to maintain absolute sensitivities of
     opical detectors, provide greater variation of correlation and increasing
     precision with which optimum correlation can be determined, especially
     when either the measured values, or known values are subject to error.
     0/3
F:
     EPI
FA
     AB; GI
     EPI: 301-A03A; 303-E04; T01-3048
Mr.
ABEQ EP
           209646 B UPAB: 19931112
     A method of sensing or determining one or more properties or the identity
     of a sample in which electromagnetic radiation is subject to optical
     interference, absorption or scatter, the method comprising the steps of:
     (a) causing electromagnetic radiation to be transmitted through, or
     reflected from said sample, said radiation including at least two
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spectrally different components so that at least one of said components is

subjected to said optical interference, abscrition or scatter and so that said components are transmitted through, or reflected from said sample by respectively different amounts; (b) measuring the transmittance or reflectance of said sample for each of said components to derive respective measured values; (c) correlating by means of either a zero dependent correlation function 'S', or a residual function 'Nres', respectively defined by: ward measured values of transmittance or reflectance with different known values representing or relating to either different values of a property of a known material, or different values which are characteristic of different known materials; and (d) selecting the known values having an optimum correlation with said measured values, the selected known values representing the property, or the identity of the sample which is sensed or to be determined.

ABEO 00 4952061 A UPAB: 19930923

Sets of measured values of the intensity of electromagnetic radiation, which has been subject to optical interference, absorption or scatter by a sample, are correlated with different sets of entwo values derived from either model of the optical properties of the sample, or from an analogue tearmique, correlation is by means of either a zero dependent correlation function, or a normalized residual function.

Both functions are unarfected by gain factors, avoiding any need to determine and to maintain absolute sensitivities of optical detectors, provide greater variation of correlation than with techniques employing a conventional correlation coefficient.

ADVANTAGE - Increased precision with which the optimum correlation can be determined, when either the measured value, or known values are subject to error. Reduced conjuting time.

L110 ANSWER 18 OF BUILDERN OF BUILDING DERWENT 1988-316508 [48] WPIN 1988-316511 [45]; 1988-316919 [4:] CF. DMN M1988-139993 Examination appts, for measuring exyperation of blood - monitors T. variations in fitting position of illumination side fixture by detecting light reflected from object. DC F31 **s03** S05 V07 HAKAMATA, N; OZAKI, T; SUZUKI, S; YASI, S 111 (HAMM) HAMAMATSU PHOTONICS KK PACYC EP 090070 A 19881109 (198845) * EM F'I1...p R: DE GB A 19900013 (199015) 11pUB 4901038 101 100000011 (109121) 111 11p 7.61E105-33 EP 136171 F: DE GB EP 190171 1 pA:1B:05-03 B1 199-0714 (1993) 5.11 A: DE GB BE 480075 B1 19930014 (19953F) E111 : · p -A61B005-05A: DE GB DE 3882272 $\mathcal{J}_{A}(\hat{j},\hat{j},\hat{k}) \triangleq \hat{j} \cdot \hat$ G 19950919 (1995%4) 7892274 $A \in (1 + 0) \cap (-0)$ 4 199:0819 (1999:4) $\mathbb{D}\mathbb{E}$ ADT EF 190.71 A EP 1988-304130 1988-355.6; US 4301.38 A US 1983-188012 19380502; EP 190271 B1 EP 1988-304130 19880500; EP 190272 B1 EP 1988-304130 1987:500; EP 240275 B1 EP 1468-704153 1477-0500; DE 3380270 3 DE 1988-3882272 19880506, EP 1988-50415 (1988-505 0); DE 3882274 G DE 1985-38-2274 19880506, EP 1998-394135 1968 0500 FDT DE 3882272 G Based on EP 290.72; DE 98.2274 G Based on EP 290.75 19870508; JP 1987-67858U 19870508; JP 1987-110465 PRAI JP 1367-113461 1987950s; JP 1987-110471 1987.5813 REP GB 2061496; GB 2075668; GB 2151620; US 3593767; US 3936192; EP 123548; EP

100768; FR 2539613; GB 2054844; US 4281645

Au1B005-00; G01N021-31; G06F015-42

ICM A61B005-00

ICS G01N021-25; G01N021-31; G06F015-42

AB EP .:90272 A UPAB: 19931116

The examination appts, measures the oxygenation using near IR light of different wavelengths. The appts, has a light source controller, an illumination side fixture, a detection side fixture a transmitted light detector and a computer to control the appts, and analyse the results. The body's heartbeat period is divided into several cycles and the transmission quantities of radiation transmitted through the head or organ are accumulated at every wavelength and for every cycle.

The computer judges whether a fitting position of the illumination side fixture has been changed on the basis of the reflection light data and the output hight data.

USE - Diagnosis of cerebral tissue damage.

Dwg.O/6

FS EPI GMPI

FA AB

MC EP1: 805-E04; 805-001

ABEQ TH 4901238 A UPAB: 19980923

The examination device comprises light source for sequentially emitting electro-magnetic waves with different wavelengths, with an illumination-side fixture for making the electro-magnetic waves introduced from the light source incident on a measuring object and detects reflected electromagnetic waves from the measuring objects. A reflection light detector detects the reflected electromagnetic waves introduced from the illumination-side fixture and outputting reflection right data. An output light detector detects emitted electromagnetic waves from the light source and outputting output light data.

A computer system receives the reflection light data from the reflection light detector and output light data from the output light detector and judges whether a fitting position of the illumination-side fixture has been changed on the basis of the reflection light data and the putput light data.

 -0.0Ξ - For measuring the oxygenation in an object with electromagnetic wave transmission spectrophotometry.

An examination apparatus (1) for measuring the oxygenation in an object (43) with electromagnetic wave transmission spectrophotometry, comprising; light source means (LD1-LD4) for sequentially emitting electromagnetic radiation of different wavelength; an illumination-side fixture (51) for contacting the electromagnetic radiation generated by the light source means (LD1-LD4) with an object (40); a receiving-side fixture (51) for receiving electrimagnetic radiation transmitted through the object (40); a transmitted light detection device (54) for detecting electromagnetic rudiation received by the receiving -side finture (fir) and a simputer system (6) for controlling the light source means (LII-LD4) and the transmitted light detecting device (50) and for analysing the output of the transmitted light detecting device (58) to determine the oxygenation of the object (45), characterised in that the illumination-side fixture (51) is arranged to receive electromagnetic waves reflected from the object (40); in that the examination apparatus also includes; reflected light detection means (4) for detecting the reflected electromagnetic radiation introduced from the illumination-side fixture (51) and outputting reflected light data; and output light detection means (13) for detecting electrimagnetic radiation emitted by the light source means (LD1-L14) and outputting output light data; and in that the computer system (6) receives the reflection light data from the reflection light detection means (4) and the output light data from the output light detection means (13), and determines whether a fitting position of the illumination-side fixture (52) changes on the basis of the reflection light data and the output light data. Ewg. 1/6

ABEQ EP 290275 B UPAB: 19931116

An examination apparatus for measuring the oxygenation of an object by electromagnetic radiation transmission spectrophotometry, comprising; light source means (LD1-LD4, 30) for sequentially emitting electromagnetic radiation at a number of different wavelengths; an illumination-side fixture (32) for applying the electromagnetic radiation emitted by the light source means (LD1-LD4) to an optect (40); transmitted light detection means (54) for notesting electrimagnetic radiation transmitted through the object (40) and outputting transmission light data; a detection-side fixture (34) for receiving electromagnetic radiation transmitted through the object (40) and crupling it to the transmitted light detection means (54); and, a computer system (56) for controlling the light source means (LCI-LG4, %) and the transmitted light detection means (54), receiving the transmission light data, and calculating the oxygenation in the object (40); characterised in that the illumination-wide fixture (3.) is equipped with a first indication means (35) for indicating if electromagnetic radiation is being emitted from the light source means (LD1-LD4); and the detection-side fixture (34) is equipped with a second indication means (55) for indicating if the transmitted light detection means (54) is in its operating condition; and, shapes and/or colours of the illumination-side fixture (32) and the detection-side fixture (54) are different to one another. Dwg.2/11

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L110 ANSWER 19 OF IC WPIM (C) 3003 THOMSON DERMENT
     1 487-137555 [05]
                          WPIE
DNN N1987-103088
     Single optical fibre transditor driving and measuring circuit - has
Τï
     bi-directional couplers recording signal intensities and transmitting
     pulsing energy to wavelength multiplexer-demultiplexer.
DC:
     P31 803 805 V07
III
     MERJCH, 3 E
FA
     (BECT) BECTON DICKINSON CO; (DESE-N) DESERET MEDICAL
     INC
CEC
    1 4
     EP 121555
                   A 198705.13 (199305) * EN
F'I
                                                   300
         F: FE OH DE ES FR GR IT LI NL SE
     UD 493m679 A 199006Le (199008)
     CA 1.80231 C 19910402 (199118)
         12355 B 199108/F (179135
R: BE CH EE FR GB IT LI LU NL
     EP 212558
     DE 3681117 G 19911002 (199141)
     FO 1000613L
                   T3 19920416 (199206)
                                                         G01DE21-31
                                                                        ·< -- --
                 Ti rasivel (1.
A. 19870814 (1999)44.
     AU 3665077
ADT EP 222555 A EP 1986-308436 1986-1025; US 4986679 A US 1985-797299 19851112; RE 19813 PR RE 1986-308436 1986-308436
         orinia, ma we 1986-208426 19861329
FDT EARLY 0132 TE Based on EP . 2. 153
PRAI UD 1985-797299 1985111.
REP As...: #13; No-SR.Pub; UU Es474xd; US 4036910; US 4114604; US 44444498
I()
     ICH G01N021-31
     ICS A61B005-00; G01N033-41
           -222585 A UPAB: 19950921
AB
     The system has an energy source connected to a power supply for emitting
     bursts of energy at a predotermined relour frequency. Another energy
     source is connected to the power supply for emitting bursts of energy at another predetermined colour frequency. A wavelength division
     multiplexer/demultiplexer is associated with the two energy sources for
     receipt of the bursts of energy, and for combination and further
     transmission.
           The discrete colour frequencies are maintained and the reflective
```

energies are separated into individual channels of each frequency upon returns of the pursts of energy. A fibre optic device and delay unit are

USE/ADVANTAGE - For in vivo measurement of blood physiological

arranged for receipt of the combined bursts of energy.

parameters. Enhances strong returns signal. 1/1

FS EFI GMPI

FAI.F.

EFI: 303-E04A9; S05-C(1; S05-D01X; V07-K04; V07-N M:C

ABEO EP 223555 B UPAB: 19930922

The system has an energy source connected to a power supply for emitting bursts of energy at a predetermined colour frequency. Another energy source is connected to the power supply for emitting bursts of energy at another predetermined colour frequency. A wavelength division multiplemer/demultiplemer is associated with the two energy sources for receipt of the bursts of energy, and for combination and further transmission.

The discrete polour frequencies are maintained and the reflective energies are separated into individual channels of each frequency upon returns of the bursts of energy. A fibre optic device and delay unit are arranged for receipt of the combined pursts of energy.

USE/ADVANTAGE - For in vivo measurement of blood physiological parameters. Enhances strong resurns signal.

1/1

4936679 A UPAB: 19930922 AREO US

> The optical fibre transducer system has an energy generator for transmitting pulsing energy at various frequencies to businectional couplers for each frequency. The couplers record the intensity and further transmit the pulsing energy to a wavelength multiplexer demultiplexer. The wavelength multiplexor/demultiplexer combines the supplies into a single output for an optic fibre which includes an optical delay sufficient to time deparate the pulsing waves of energy. Reflected energy is transmitted back through the same wavelength multiplexer demultiplexer, bidirectional compler so that the recorder intensity of transmission and reflectance are comparable with system influence.

> A method is also shown for use of an optical fibre system including the components set forth and the system requires the generation and combination of the various frequencies of energy in a multiplexer/demultiplexer, the delaw for time separation and the detection in a bigirection couplor of transmitted and reflected energy. MSE - Catheter instrument.

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LIIO ANGWER 20 OF 20 WPIN (C) 1002 THOMNON DERWENT
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AH - 1981-64/00D: [86] WPIX

Investigating unknown substance - by comparing spectral peak table for Τ'nΙ substance with library of chemical structural units.

[1(~ JC1 803

I1: CARTER, H V; COATES, J F; FORE, M A; HANNAH, R W; SAVITERY, A

(FERE: FERFIN-ELMER CORE F 7-.

 $\subseteq \Gamma \subseteq \Gamma$

F-1 GB U670U75 A 19811 + 15 1981360 * 17pA 198111. 6 (198149) A 19821221 (198302) B 19840201 (198405) DE 3104178

US 4365303

GB 1670235

GE L070L35 A GB 1981-3088 19810202 ALT

PRAI US 1980-119837 19800207

ICG01N021-31; G06F015-20 GE 25.76235 A UEAB: 1 4930 415 АБ

Apprts, for determining the nature of an unknown substance can enter a spectral peak table for the substance into computing appts, and adjust it to a first preselected standardised format and compare it with a first library of enemical structural units in a memory. The possible units most closely corresponding to the unknown substance are listed.

The peak table is readjusted to a second preselected standardised format. A file is formed for the unknown substance including its readjusted peak table and data of the list of possible chemical structural units is compared with files in a second library each contg. data for a

known substance and including its readjusted peak table and data of its chemical structural unit. A list of known substances most closely corresponding to the unknown substance is presented.

FS CPI EPI

FA AB

MC CFI: J04-E01A

EPI: SCH-E04A9; SUR-E09X

ABEQ GE 2070135 B UPAB: 19930915

Appts, for determining the nature of an unknown substance can enter a spectral peak tuble for the substance into computing appts, and adjust it to a first presolested standardised format and compare it with a first library of chemical structural units in a memory. The possible units most closely corresponding to the unknown substance are listed.

The peak table is readjusted to a second preselected standardised format. A file is formed for the unknown substance including its readjusted peak table and data of the list of possible chemical structural units is compared with files in a second library each contg. data for a known substance and including its readjusted peak table and data of its chemical structural unit. A list of known substances most closely corresponding to the unknown substance is presented.

$= \cdot d his$

(FILE 'HOME' ENTERED AT 06:39:28 ON 08 JUL 2002) SET COST OFF

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FILE 'HCAPLUS' ENTERED AT 06:39:41 ON 08 JUL 2002
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```
E WILES T /AU
L1
                1 S E6
                  E TUFNER D/AU
L.:
             109 S E3, E15
               30 S EDS, E44
L_{-}
              14 3 E60, E61
               1 3 E41
                  E O CONNEL M/AU
               1 S E6
Li
                  E O CONNELL MYAU
               3 4 8 E1, NA
              45 3 E51,E58,E61
                 E PARMIGIANI G/AU
              10 S E5
LH
                  E CLYDE M/AU
L10
               . S E4, E(, E7
             987 S (RECTON? (L) DICKIN?) /PA, GS
T.11
                  E MATHEMATIC/CT
                 E E : ALI.
L12
            29640 € E1
                 E E. ·ALL
             491 S E3, E5
            5666 S E2
L15
           81263 S ED+NT
          171388 S MATH?
L16
L17
          19037 S L16 AND L12-L15
Lla
          239020 S L11-L17
                  E TURRIDITY/OT
                  E E3+ALL
L1 3
            1609 S Ex
           36976 S TUFFIDE
L.:0
            26\%2 S E6+MT OR E9+NT
L_{-}1
             -6. ↔ S E1. ·· NT
L.2
L \angle 3
            1860 3 E4, 5I
```

E E5+ALL

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L.7.4
            4453 S E5
                  E REDOM/CT
                  E E32+ALL
            790 S E-,E9
LT:5
            5188 S E7
Lib
LJ
           1830# S E7.NT
                  E ELM-ALL
L^{r_{1}\otimes}
          318745 S E2+NT
          187396 S EGS+NT OR E70+NT OR E71+NT OR E72+NT
L. .:
          10078% S REDOM
L:0
L- 1
             540 S L18 AND L19-L24
L \cdot \Box
            5241 S L18 AND L25-L30
L 5 5
              - 6 S L31 AND L32
            5775 S L31, L31
L:4
              GG S L34 AND (WAVELENG? OR WAVE LENG?)
L
                  E LIGHT/CT
                  E E - ALL
Lit
             704 S L1: AND E8:NT
\mathbb{L}\,\mathbb{R}^+
            900A S Lie AND (E33+NT OF E23+NT OF E25+NT OR E26+NT OR E27+NT OF E2
             7.0 S LIT AND (E40+NT OF E47+NT OR E48+NT OR E49+NT OR E50+NT OR E5
L_{-}: \vdash
            6120 S LIF AND LIGHT?
L \beta \beta
            2004 S Lit AMP (WAVELENG? OF WAVE LENG?)
Lil
              26 S Lle AND WL
1.44
              MIX S LEW-LAT AND LET
L4..
             2501 S LEW-L41 AND L32
L4
               4 S L4. AND L43
L.; 4
                  E SCREENING/CT
                  E EFFALL
                  E DEGG SCREENING/CT
                  E = E + ALL
L.:
           18485 S EL, El + NT
            45 35 S E5+NT
L: \psi
L : 7
                1 8 L41, L47 AND L45, L46
                  E APPARATUS/CT
L4^{\frac{1}{4}}
                O S L43, L43 AND E3
                . S L43, L43 AND E3/CW
Lin
                  E MEASURING APPARATUS/CT
                  E EstAM
                4 3 Lit AND E4,E5
Lin
             Build S Life AND ESHIT
T . .
                  E Ebel +ALL
              s'i s bis AND E3,E2+NT
                  E ESSHALL
L'·
              85 S LIS AND ED-NT
L:.:
              M : C DI. ZEE EE-NT
\Gamma_{i}:
             1287 S L18 AND EXHIT
            3×75 S L49-L55
L56
\Gamma
             L'. ~
              d/ 3 Lie AMD Lie
              5.1 S 157,15%
Lt.
            10MR S LIN AND GOIN/IC, ICM, ICS
Len
Lel
              -4 S Le0 AM L34
              JA S LOI AME LB5-L50
Litti
               34 3 LGT-LG.
Lbs
              . 5 S L59 AND (9 OR 10 OR 4)/SC,SX
Line
               18 3 15: NOT Les, 164
Lon
                  SEL DN AN 3 6
                . S El-E6
Lim
L\tilde{\phi}^{-}
               4 # 8 20 4, 204, 100
               31 S LI-LII AND LIB
L_{\mathcal{D}^{\pm}}
               + S Lo3 AND L13-L67
L6 +
                E O CONNELL M. AU
```

L7J

50 S E3, E4

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19 S E51,E52,E60
L71
L711
               1 S L18 AND L70, L71
               8 S L68, L72 AND L19-L67
L73
                 SEL LN AN 5-8
               4 S L7: NOT EL-E12
L74
              38 S L67 AND (PEK=20006531 OF FEEK=20000531 OR AD<=20000531)
L75
              41 S L74,175
L76
L77
              41 S L76 AND L1-L76
                 SEL DN AN L77 4 6 10 11 16 17 21 24-28 33 34 38 39 41
              24 S L77 NOT E13-E63
L7::
              14 S L78 AND (GFOWT OF CONCENTER OF LIGHT? OF WAVELENG? OF WAVE LE
L7:
     FILE 'HCAPLUS' ENTERED AT 07:32:51 OF 08 JUL 2002
             HS S LIW AND GOINOHI-SI'IC, ICM, ICM
L80
              34 S L80 NOT L79
L \in I
              20 S L81 AND (FT) = 20000531 OF FED = 20000531 OF ADK = 20000531)
L::
              20 S L82 AMO L1-180
Lis
                 SEL 183 DN AN D 4 5 6 12 13 14 16 17 20
L54
              10 S L83 NOT E64-E93
              10 S L84 AMD (GROW? OR CONCENTE? OF LIGHT? OF WAVELENG? OF WAVE LE
L35
     FILE 'WPIM' ENTERED AT 07:41:24 ON 06 JUL 2002
                 E WILES T/AU
L \oplus G
               1 S E4
                 E O CONNELL M/AU
L87
              10 S E3, E4
                 E O CONNEL M/AU
               1 S E3
L88
                 E OCCUMELL M/AU
               a S E3,E4
Lab
                 E PARMIGIANI F'AU
               2 S E3
L90
                 E TUFNER D/AU
              55 S E3,E13,E14
L91
                 E CLYDE M/AU
               _ S E3
L92
                 E BECT/PA
            2073 S (BECTO(L)DICKE)/PA
L93
                 E BECT/PACO
                 E E3+ALL
L94
            2070 S E1
            1386 S G01N021-31/10, ICM, IC3
L95
               6 S L95 AND 012M001-34/IC, ICM, ICS
L96
             323 S 195 AND GOINGBE/IC, ICM, ICS
L97
               0 a 600P/10,10M,100 AMD 107
L90
L99
               4 S LSe-L94 AND L95
L1:)(+
               9 S L98, L99 AND L96, L97
L101
               4 S L100 AND GUIN/IC, ICM, ICU
                 SEL DU AU 4-8
               4 S L101 NOT E1-E13
L1:02
L100:
               7 S LB6, LB--L100 NOT L101
                 SEL DN AN 1
               1 S L103 AND E14-E15
L1:04
               5 S L102, L104
L1:05
               6 S L99, L105
L10v
              25 S L95 AND GU6F/IC, ICM, ICS
L107
              17 S L107 NOT L98-L106
L100
              23 S L100, L108 AND L86-L108
L104
              20 S S03/DC AND 1109
L11:
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FILE 'WPIK' ENTERED AT 03:01:46 ON 08 JUL 2002